PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification. 6:

C12N 15/82, 9/12, C07K 14/415, A01H
5/00

(11) International Publication Number: WO 97/43427

(43) International Publication Date: 20 November 1997 (20.11.97)

(21) International Application Number: PCT/EP97/02443 (81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR,

13 May 1997 (13.05.97)

(30) Priority Data: 9610044.1 14

(22) International Filing Date:

14 May 1996 (14,05.96)

GB

(71) Applicant (for all designated States except US): NOVARTIS AG [CH/CH]; Scharzwaldallee 215, CH-4058 Basel (CH).

(72) Inventors; and

(75) Inventors/Applicants (for US only): DE VRIES, Sape, Cornelis [NL/NL]; Roghorst 192, NL-6708 KS Wageningen (NL). SCHMIDT, Eduard, Daniel, Leendert [NL/NL]; Callunastraat 25, NL-6813 ET Arnheim (NL). VAN HOLST, Gerrit, Jan [NL/NL]; De Gouw 8, NL-1602 DN Enkhuizen (NL). HECHT, Valerie, France, Gabrielle [NL/NL]; Kees Muldenweg 25, NL-6707 HA Wageningen (NL).

(74) Agent: ROTH, Bernhard, M.; Novartis AG, Patent- und Markenabteilung, Klybeckstrasse 141, CH-4002 Basel (CH).

(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, ARIPO patent (GH, KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: PRODUCTION OF APOMICTIC SEED

(57) Abstract

The present invention provides, inter alia, a method of producing apomictic seeds comprising the steps of: (i) transforming plant material with a nucleotide sequence encoding a protein the presence of which in a cell, or membrane thereof, renders said cell embryogenic, (ii) regenerating the thus transformed material into plants, or carpel containing parts thereof, and (iii) expressing the sequence in the vicinity of the embryo sac. The protein may be a leucine repeat rich receptor kinase which preferably is modified to the extent that the ligand binding domain is deleted or functionally inactivated.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	Fl	Finland	LT	Lithuania	SK	Slovakia
ΑT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑÜ	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
A7.	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
вв	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	MI.	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	1T	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ.	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Gеппалу .	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		•
EE	Estonia	LR	Liberia	SG	Singapore		

WO 97/43427 PCT/EP97/02443

Production of Apomictic Seed

The present invention relates to the production of genetically transformed plants. In particular the invention relates *inter alia* to a process for inducing apomixis, to the apomictic seeds which result from the process, and to the plants and progeny thereof which result from the germination of such seeds.

Apomixis, which is vegetative (non-sexual) reproduction through seeds, is a genetically controlled reproductive mechanism found in some polyploid non-cultivated species. The process is classified as gametophytic or non-gametophytic. In gametophytic apomixis - of which there are two types (apospory and diplospory), multiple embryo sacs which typically lack antipodal nuclei are formed, or else megasporogenesis in the embryo sac takes place. In adventitious embryony (non-gametophytic apomixis), a somatic embryo develops directly from the cells of the embryo sac, ovary wall or integuments. In adventitious embryony, somatic embryos from surrounding cells invade the sexual ovary, one of the somatic embryos out-competes the other somatic embryos and the sexual embryo and utilizes the produced endosperm.

Were apomixis to be a controllable and reproducible phenomenon it would provide many advantages in plant improvement and cultivar development in the case that sexual plants are available as crosses with the apomictic plant.

For example, apomixis would provide for true-breeding, seed propagated hybrids. Moreover, apomixis could shorten and simplify the breeding process so that selfing and progeny testing to produce and/or stabilize a desirable gene combination could be eliminated. Apomixis would provide for the use as cultivars of genotypes with unique gene combinations since apomictic genotypes breed true irrespective of heterozygosity. Genes or groups of genes could thus be "pyramided and "fixed" in super genotypes. Every superior apomictic genotype from a sexual-apomictic cross would have the potential to be a cultivar. Apomixis would allow plant breeders to develop cultivars with specific stable traits for such characters as height, seed and forage quality and maturity. Breeders would not be limited in their commercial production of hybrids by (i) a cytoplasmic-nuclear interaction to produce male sterile female parents or (ii) the fertility restoring

capacity of a pollinator. Almost all cross-compatible germplasm could be a potential parent to produce apomictic hybrids.

Finally, apomixis would simplify commercial hybrid seed production. In particular, (i) the need for physical isolation of commercial hybrid production fields would be eliminated; (ii) all available land could be used to increase hybrid seed instead of dividing space between pollinators and male sterile lines; and (iii) the need to maintain parental line seed stocks would be eliminated.

The potential benefits to accrue from the production of seed *via* apomixis are presently unrealized, to a large extent because of the problem of engineering apomictic capacity into plants of interest. The present invention provides a solution to that problem in that it provides the means for obtaining plants which exhibit the adventitious embryony type of apomixis.

According to the present invention there is provided a method of producing apomictic seeds comprising the steps of:

- (i) transforming plant material with a nucleotide sequence encoding a protein the presence of which in an active form in a cell, or membrane thereof, renders said cell embryogenic,
- (ii) regenerating the thus transformed material into plants, or carpel-containing parts thereof, and
- (iii) expressing the sequence in the vicinity of the embryo sac.

By "vicinity of the embryo sac" is meant in one or more of the following: carpel, integuments, ovule, ovule premordium, ovary wall, chalaza, nucellus, funicle and placenta. The skilled man will recognize that the term "integuments" also includes those tissues, such as endothelium, which are derived therefrom. By "embryogenic" is meant the capability of cells to develop into an embryo under permissive conditions. It will be appreciated that the term "in an active form" includes proteins which are truncated or otherwise mutated with the proviso that they initiate or amplify embryogenesis whether or not in doing this they interact with the signal transduction components that they otherwise would in the tissues in which they are normally present.

The term "plant material" includes protoplasts, isolated plant cells (such as stomatal guard cells) possessing a cell wall, pollen, whole tissues such as emerged radicle, stem, leaf, petal,

WO 97/43427 PCT/EP97/02443

hypocotyl section, apical meristem, ovaries, zygotic embryo per se, roots, vascular bundle, pericycle, anther filament, somatic embryos and the like.

A further embodiment of the invention relates to a DNA molecule comprising a nucleotide sequence encoding a protein the presence of which in an active form in a cell, or membrane thereof, renders said cell embryogenic.

The said nucleotide sequence may be introduced into the plant material, *inter alia*, *via* a bacterial or viral vector, by micro-injection, by co-incubation of the plant material and sequence in the presence of a high molecular weight glycol or by coating of the sequence onto the surface of a biologically inert particle which is then introduced into the material.

Expression of the sequence may yield a protein kinase capable of spanning a plant cell membrane. Typically the kinase may be a leucine rich repeat receptor like kinase which has the capacity to auto-phosphorylate. The skilled man will recognize what is meant by the term "leucine rich repeat receptor like kinase". Examples of such proteins include Arabidopsis RLK5 (Walker, 1993), Arabidopsis RPS2 (Bent et al. 1994), Tomato CF-9 gene product (Jones et al. 1994), Tomato N (Whitham et al. 1994), Petunia PRK1 (Mu et al. 1994), the product of the Drosophila Toll gene (Hashimoto et al. 1988), the protein kinase encoded by the rice OsPK10 gene (Zhao et al. 1994), the translation product of the rice EST clone ric2976 and the product of the Drosophila Pelle gene (Shelton and Wasserman, 1993). Still further examples of such proteins include the TMK1, Clavatal, Erecta, and TMKL1 gene products from Arabidopsis, the Flightless-1 gene product from Drosophila, the TrkC gene product from pig, the rat LhCG receptor and FSH receptor, the dog TSH receptor, and the human Trk receptor kinase. The protein may comprise a ligand binding domain, a proline box, a transmembrane domain, a kinase domain and a protein binding domain. In many receptor kinases the extracellular (ligand binding) domain serves as an inhibitor of the kinase domain in the ligand-free state. This arrest is removed after binding of the ligand. Accordingly, in one embodiment of the invention the protein either lacks a ligand binding domain or the domain is functionally inactivated so that the kinase domain can be constitutively active in the absence of an activating signal (ligand). Whether or not the protein possesses a ligand binding domain - functional or otherwise, once expressed and incorporated into the plant cell membrane the protein binding domain is preferably located intra-cellularly.

In a preferred embodiment of the method, the said sequence further encodes a cell membrane targeting sequence. The sequence may be that which is depicted in SEQ ID Nos. 1, 2, 20, or 32, or it may be similar in that it is complementary to a sequence which hybridizes under stringent conditions with the said sequences and which encodes a membrane bound protein having kinase activity. By "similar" is meant a sequence which is complementary to a test sequence which is capable of hybridizing to the inventive sequence. When the test and inventive sequences are double stranded the nucleic acid constituting the test sequence preferably has a TM within 20°C of that of the inventive sequence. In the case that the test and inventive sequences are mixed together and denatured simultaneously, the TM values of the sequences are preferably within 10°C of each other. More preferably the hybridization is performed under stringent conditions, with either the test or inventive DNA preferably being supported. Thus either a denatured test or inventive sequence is preferably first bound to a support and hybridization is effected for a specified period of time at a temperature of between 50 and 70°C in double strength citrate buffered saline (SSC) containing 0.1%SDS followed by rinsing of the support at the same temperature but with a buffer having a reduced SSC concentration. Depending upon the degree of stringency required, and thus the degree of similarity of the sequences, at a particular temperature, - such as 60°C, for example - such reduced concentration buffers are typically single strength SSC containing 0.1%SDS, half strength SSC containing 0.1%SDS and one tenth strength SSC containing 0.1%SDS. Sequences having the greatest degree of similarity are those the hybridization of which is least affected by washing in buffers of reduced concentration. It is most preferred that the test and inventive sequences are so similar that the hybridization between them is substantially unaffected by washing or incubation in one tenth strength sodium citrate buffer containing 0.1%SDS.

Accordingly, further comprised by the present invention is a DNA sequence as depicted in SEQ ID NOS: 22, 24, 26, 28 and 30 or a sequence which is complementary to one which hybridizes under stringent conditions with the said sequences and which encodes a membrane bound protein having kinase activity.

The sequence may be modified in that known mRNA instability motifs or polyadenylation signals may be removed and/or codons which are preferred by the plant into which the sequence is to be inserted may be used so that expression of the thus modified sequence in the said plant may yield substantially similar protein to that obtained by expression of the unmodified sequence in the organism in which the protein is endogenous.

In order to obtain expression of the sequence in the regenerated plant (and in particular the carpel thereof) in a tissue specific manner the sequence is preferably under expression control of an inducible or developmentally regulated promoter, typically one of the following: a promoter which regulates expression of SERK genes *in planta*, the *Arabidopsis* ANT gene promoter, the

which regulates expression of SERK genes *in planta*, the *Arabidopsis* ANT gene promoter, the promoter of the O126 gene from *Phalaenopsis*, the carrot chitinase DcEP3-1 gene promoter, the *Arabidopsis* AtChitlV gene promoter, the *Arabidopsis* LTP-1 gene promoter, the *Arabidopsis* bellogene promoter, the petunia fbp-7 gene promoter, the *Arabidopsis* AtDMC1 promoter, the pTA7001 inducible promoter. The DcEP3-1 gene is expressed transiently during inner integument degradation and later in cells that line the inner part of the developing endosperm. The AtChilV gene is transiently expressed in the micropylar endosperm up to cellularisation. The LTP-1 promoter is active in the epidermis of the developing nucellus, both integuments, seed coat and early embryo. The bel-1 gene is expressed in the developing inner integument and the fbb-7 promoter is active during embryo sac development. The *Arabidopsis* ANT gene is expressed during integument development, and the O126 gene from *Phalaenopsis* is expressed in the mature ovule.

It is most preferred that the sequence is expressed in the somatic cells of the embryo sac, ovary wall, nucellus, or integuments.

The endosperm within the apomictic seed results from fusion of polar nuclei within the embryo sac with a pollen-derived male gamete nucleus. It is preferred that the sequence encoding the protein is expressed prior to fusion of the polar nuclei with the male gamete nucleus.

The invention further includes a DNA, but preferably a recombinant DNA, comprising a sequence encoding a protein the presence of which in an active form in a cell, or membrane thereof, renders said cell embryogenic. Preferred is a DNA encoding a protein which is a leucine nich repeat receptor like kinase and comprises a ligand binding domain, a proline box, a transmembrane domain, a kinase domain and a protein binding domain, the ligand binding domain optionally being absent or functionally inactive.

In particular, the invention embodies a DNA comprising a DNA sequence encoding a N-terminal protein fragment having the following amino acid sequence: Gin Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val Thr Cys Asn.

WO 97/43427 PCT/EP97/02443

A specific embodiment of the invention relates to a DNA comprising a DNA sequence encoding a protein having the sequence depicted in SEQ ID Nos. 3 or 21, or a protein substantially similar thereto which is capable of being membrane bound and which has kinase activity. By substantially similar is meant a pure protein having an amino acid sequence which is at least 90% similar to the sequence of the proteins depicted in SEQ ID No 3 below. In the context of the present invention, two amino acid sequences with at least 90% similarity to each other have at least 90% identical or conservatively replaced amino acid residues in a like position when aligned optimally allowing for up to 8 gaps with the *proviso* that in respect of each gap a total not more than 4 amino acid residues is affected. For the purpose of the present invention conservative replacements may be made between amino acids within the following groups:

- (i) Serine and Threonine;
- (ii) Glutamic acid and Aspartic acid;
- (iii) Arginine and Lysine;
- (iv) Asparagine and Glutamine;
- (v) Isoleucine, Leucine, Valine and Methionine;
- (vi) Phenylalanine, Tyrosine and Tryptophan
- (vii) Alanine and Glycine

In addition, non-conservative replacements may also occur at a low frequency. Accordingly, the invention futher embodies a DNA comprising a DNA sequence encoding a N-terminal protein fragment having the following amino acid sequence: Val Xaa Gin Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val Thr Cys Asn, with Xaa being a variable amino acid, but preferably Leu or Val.

Especially preferred within the scope of the invention is a DNA comprising a DNA sequence encoding a N-terminal protein fragment having the following amino acid sequence: Val Xaa Gin Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val Thr Cys Asn Xab Xac Xad Xae Val Xaf Arg Val Asp Leu Giy Asn Xag Xah Leu Ser Giy His Leu Xai Pro Giu Leu Giy Xaj Leu Xak Xal Leu Gin, with Xaa to Xak representing variable amino acids, but preferably

Xaa = Leu or Val

Xab = AsnorGin

Xac = Gluor Aspor His

Xad = AsnorHis

Xae = Seror Argor Gin

WO 97/43427 PCT/EP97/02443

Xaf = leorThr

Xag = Ala or Ser

Xah = Glu or Asn

Xai = Valor Ala

Xai= Valor Lys

Xak=LysorGiu

Xal = Asn or His

It is preferred that the DNA further encodes a cell membrane targeting sequence, and that the protein encoding region is under expression control of a developmentally regulated or inducible promoter, such as, for example, a promoter which regulates expression of SERK genes in planta, the carrot chitinase DcEP3-1 gene promoter, the Arabidopsis AtChittV gene promoter, the Arabidopsis LTP-1 gene promoter, the Arabidopsis bel-1 gene promoter, the petunia fbp-7 gene promoter, the Arabidopsis ANT gene promoter, or the promoter of the O126 gene from Phalaenopsis; the Arabidopsis AtDMC1 promoter, or the pTA7001 inducible promoter.

Particularly preferred embodiments of the said DNA include those depicted in SEQ ID Nos. 1, 2, 20 or 32, or those which are complementary to one which hybridizes under stringent conditions with the said sequences and which encode a membrane bound protein having kinase activity. As indicated above, the DNA may be modified in that known mRNA instability motifs or polyadenylation signals may be removed and/or codons which are preferred by the plant into which the DNA is to be inserted may be used so that expression of the thus modified DNA in the said plant may yield substantially similar protein to that obtained by expression of the unmodified DNA in the organism in which the protein is endogenous.

The invention still further includes a vector which contains DNA as indicated in the three immediately preceding paragraphs, plants transformed with the recombinant DNA or vector, and the progeny of such plants which contain the DNA stably incorporated, and/or the apomictic seeds of such plants or such progeny.

The recombinant DNA molecules of the invention can be introduced into the plant cell in a number of art-recognized ways. Those skilled in the art will appreciate that the choice of method might depend on the type of plant, i.e. monocot or dicot, targeted for transformation. Suitable methods of transforming plant cells include microinjection

(Crossway et al., BioTechniques 4:320-334 (1986)), electroporation (Riggs et al, Proc. Natl. Acad. Sci. USA 83:5602-5606 (1986), Agrobacterium mediated transformation (Hinchee et al., Biotechnology 6:915-921 (1988)), direct gene transfer (Paszkowski et al., EMBO J. 3:2717-2722 (1984)), ballistic particle acceleration using devices available from Agracetus, Inc., Madison, Wisconsin and Dupont, Inc., Wilmington, Delaware (see, for example. Sanford et al., U.S. Patent 4,945,050; and McCabe et al., Biotechnology 6:923-926 (1988)), and protoplast transformation/regeneration methods (see U.S. Patent No. 5,350,689 issued Sept. 27, 1994 to Ciba-Geigy Corp.). Also see, Weissinger et al., Annual Rev. Genet. 22:421-477 (1988); Sanford et al., Particulate Science and (1987)(onion); Christou et al., Plant Physiol. 87:671-674 Technology 5:27-37 (1988)(soybean); McCabe et al., Bio/Technology 6:923-926 (1988)(soybean); Datta et al., Bio/Technology 8:736-740 (1990)(rice); Klein et al., Proc. Natl. Acad. Sci. USA, 85:4305-4309 (1988)(maize); Klein et al., Bio/Technology 6:559-563 (1988)(maize); Klein et al., Plant Physiol. 91:440-444 (1988)(maize); Fromm et al., Bio/Technology 8:833-839 (1990); and Gordon-Kamm et al., Plant Cell 2:603-618 (1990)(maize).

Comprised within the scope of the present invention are transgenic plants, in particular transgenic fertile plants transformed by means of the aforedescribed processes and their asexual and/or sexual progeny, which still contain the DNA stably incorporated, and/or the apomictic seeds of such plants or such progeny.

The transgenic plant according to the invention may be a dicotyledonous or a monocotyledonous plant. Such plants include field crops, vegetables and fruits including tomato, pepper, melon, lettuce, cauliflower, broccoli, cabbage, brussels sprout, sugar beet, com, sweetcom, onion, carrot, leek, cucumber, tobacco, alfalfa, aubergine, beet, broad bean, celery, chicory, cow pea, endive, gourd, groundnut, papaya, pea, peanut, pineapple, potato, safflower, snap bean, soybean, spinach, squashes, sunflower, sorghum, water-melon, and the like; and omamental crops including Impatiens, Begonia, Petunia, Pelargonium, Viola, Cyclamen, Verbena, Vinca, Tagetes, Primula, Saint Paulia, Ageratum, Amaranthus, Anthirmhinum, Aquilegia, Chrysanthemum, Cineraria, Clover, Cosmo, Cowpea, Dahlia, Datura, Delphinium, Gerbera, Gladiolus, Gloxinia, Hippeastrum, Mesembryanthemum, Salpiglossis, Zinnia, and the like. In a preferred embodiment, the DNA is expressed in "seed crops" such as com, sweet com and peas etc. in such a way that the apomictic seed which results from such expr ssion is not physically mutated or otherwise damaged in comparison with seed

from untransformed like crops. Preferred are monocotyledonous plants of the Graminaceae family involving <u>Lolium</u>, <u>Zea</u>, <u>Triticum</u>, <u>Triticale</u>, <u>Sorghum</u>, <u>Saccharum</u>, <u>Bromus</u>, <u>Orvzae</u>, <u>Avena</u>, <u>Hordeum</u>, <u>Secale</u> and <u>Setaria</u> plants.

More preferred are transgenic maize, wheat, barley, sorghum, rye, oats, turf and forage grasses, millet and rice. Especially preferred are maize, wheat, sorghum, rye, oats, turf grasses and rice.

Among the dicotyledonous plants Arabidopsis, soybean, cotton, sugar beet, sugar cane, oilseed rape, tobacco and sunflower are more preferred herein. Especially preferred are soybean, cotton, tobacco, sugar beet and oilseed rape.

The expression 'progeny' is understood to embrace both, "asexually" and "sexually" generated progeny of transgenic plants. This definition is also meant to include all mutants and variants obtainable by means of known processes, such as for example cell fusion or mutant selection and which still exhibit the characteristic properties of the initial transformed plant, together with all crossing and fusion products of the transformed plant material. This also includes progeny plants that result from a backcrossing, as long as the said progeny plants still contain the DNA according to the invention

Another object of the invention concerns the proliferation material of transgenic plants.

The proliferation material of transgenic plants is defined relative to the invention as any plant material that may be propagated sexually or asexually *in vivo* or *in vitro*. Particularly preferred within the scope of the present invention are protoplasts, cells, calli, tissues, organs, seeds, embryos, pollen, egg cells, zygotes, together with any other propagating material obtained from transgenic plants.

Parts of plants, such as for example flowers, stems, fruits, leaves, roots originating in transgenic plants or their progeny previously transformed by means of the process of the invention and therefore consisting at least in part of transgenic cells, are also an object of the present invention. Espeically preferred a apomictic seeds.

A further object of the invention is a method of producing apomictic seeds, but preferably seeds that are of the adventitious embryony type, comprising the steps of:

(i) transforming plant material with a nucleotide sequence encoding a protein the presence of which in an active form in a cell, or membrane thereof, renders said cell

embryogenic, but preferably a protein which is a protein kinase capable of spanning a plant cell membrane and capable of autophosphorylation.

- (ii) regenerating the thus transformed material into plants, or carpel-containing parts thereof, and
- (iii) expressing the sequence in the vicinity of the embryo sac.

The kinase protein being expressed by the DNA according to the invention is preferably a leucine rich repeat receptor like kinase and comprises a ligand binding domain, a proline box, a transmembrane domain, a kinase domain and a protein binding domain. In a specific embodiment of the invention, the said kinase protein may lack a functional ligand binding domain but comprises a proline box, a transmembrane domain, a kinase domain and a protein binding domain.

The genetic properties engineered into the transgenic seeds and plants described above are passed on by sexual reproduction or vegetative growth and can thus be maintained and propagated in progeny plants. Generally said maintenance and propagation make use of known agricultural methods developed to fit specific purposes such as tilling, sowing or harvesting. Specialized processes such as hydroponics or greenhouse technologies can also be applied. As the growing crop is vulnerable to attack and damages caused by insects or infections as well as to competition by weed plants, measures are undertaken to control weeds, plant diseases, insects, nematodes, and other adverse conditions to improve yield. These include mechanical measures such a tillage of the soil or removal of weeds and infected plants, as well as the application of agrochemicals such as herbicides, fungicides, gametocides, nematicides, growth regulants, ripening agents and insecticides.

Use of the advantageous genetic properties of the transgenic plants and seeds according to the invention can further be made in plant breeding which aims at the development of plants with improved properties such as tolerance of pests, herbicides, or stress, improved nutritional value, increased yield, or improved structure causing less loss from lodging or shattering. The various breeding steps are characterized by well-defined human intervention such as selecting the lines to be crossed, directing pollination of the parental lines, or selecting appropriate progeny plants. Depending on the desired properties different breeding measures are taken. The relevant techniques are well known in the art and include but are not limited to hybridization, inbreeding, backcross breeding, multiline breeding,

variety blend, interspecific hybridization, aneuploid techniques, etc. Hybridization techniques also include the sterilization of plants to yield male or female sterile plants by mechanical, chemical or biochemical means. Cross pollination of a male sterile plant with pollen of a different line assures that the genome of the male sterile but female fertile plant will uniformly obtain properties of both parental lines. Thus, the transgenic seeds and plants according to the invention can be used for the breeding of improved plant lines which for example increase the effectiveness of conventional methods such as herbicide or pestidice treatment or allow to dispense with said methods due to their modified genetic properties. Alternatively new crops with improved stress tolerance can be obtained which, due to their optimized genetic "equipment", yield harvested product of better quality than products which were not able to tolerate comparable adverse developmental conditions.

In seeds production germination quality and uniformity of seeds are essential product characteristics, whereas germination quality and uniformity of seeds harvested and sold by the farmer is not important. As it is difficult to keep a crop free from other crop and weed seeds, to control seedborne diseases, and to produce seed with good germination, fairly extensive and well-defined seed production practices have been developed by seed producers, who are experienced in the art of growing, conditioning and marketing of pure seed. Thus, it is common practice for the farmer to buy certified seed meeting specific quality standards instead of using seed harvested from his own crop. Propagation material to be used as seeds is customarily treated with a protectant coating comprising herbicides, insecticides, fungicides, bactericides, nematicides, molluscicides or mixtures thereof. Customarily used protectant coatings comprise compounds such as captan, carboxin, thiram (TMTD*), methalaxyl (Apron*), and pirimiphos-methyl (Actellic*). If desired these compounds are formulated together with further carriers, surfactants or applicationpromoting adjuvants customarily employed in the art of formulation to provide protection against damage caused by bacterial, fungal or animal pests. The protectant coatings may be applied by impregnating propagation material with a liquid formulation or by coating with a combined wet or dry formulation. Other methods of application are also possible such as treatment directed at the buds or the fruit.

It is thus a further object of the present invention to provide plant propagation material for cultivated plants, but especially plant seed that is treated with an seed protectant coating customarily used in seed treatment.

It is a further aspect of the present invention to provide new agricultural methods such as the methods examplified above which are characterized by the use of transgenic plants, transgenic plant material, or transgenic seed according to the present invention.

To breed progeny from plants transformed according to the method of the present invention, a method such as that which follows may be used: plants produced as described in the examples set forth below are grown in pots in a greenhouse or in soil, as is known in the art, and permitted to flower. Pollen is obtained from the mature stamens and used to pollinate the pistils of the same plant, sibling plants, or any desirable plant. Similarly, the pistils developing on the transformed plant may be pollinated by pollen obtained from the same plant, sibling plants, or any desirable plant. Transformed progeny obtained by this method may be distinguished from non-transformed progeny by the presence of the introduced gene(s) and/or accompanying DNA (genotype), or the phenotype conferred. The transformed progeny may similarly be selfed or crossed to other plants, as is normally done with any plant carrying a desirable trait. Similarly, tobacco or other transformed plants produced by this method may be selfed or crossed as is known in the art in order to produce progeny with desired characteristics. Similarly, other transgenic organisms produced by a combination of the methods known in the art and this invention may be bred as is known in the art in order to produce progeny with desired characteristics.

Further comprised by the invention is a method of obtaining embryogenic cells in plant material, comprising transforming the material with a recombinant DNA sequence or a vector according to the invention, expressing the sequence in the material or derivatives thereof and subjecting the said material or derivatives to a compound which acts as a ligand for the gene product of the said sequence.

The invention further relates to a method of generating somatic embryos under in vitro conditions wherein the SERK protein is overexpressed ectopically.

The invention still further includes the use of the said DNA in the manufacture of apomictic seeds, in which use the sequence is expressed in the vicinity of the embryo sac.

In a specific embodiment of the invention the SERK gene may be expressed in transgenic plants such as, for example, an *Arabidopsis* plant, under the control of plant expression signals, particularly a promoter which regulates expression of SERK genes *in planta*, but preferably a

WO 97/43427 PCT/EP97/02443

developmentally regulated or inducible promoter such as, for example, the carrot chitinase DcEP3-1 gene promoter, the *Arabidopsis* AtChitlV gene promoter, the *Arabidopsis* LTP-1 gene promoter, the *Arabidopsis* bel-1 gene promoter, the petunia fbp-7 gene promoter, the *Arabidopsis* ANT gene promoter, or the promoter of the O126 gene from *Phalaenopsis*; the *Arabidopsis* AtDMC1 promoter, or the pTA7001 inducible promoter.

The promoters of the DcEP3-1 and the AtChit IV genes may be cloned and characterized by standard procedures. The DcSERK coding sequence (SEQ ID No. 2) is cloned behind the DcEP3-1, the AtChit IV or the AtLTP-1 promoters and transformed into Arabidopsis. The ligation is performed in such a way that the promoter is operably linked to the sequence to be transcribed. This construct, which also contains known marker genes providing for selection of transformed material, is inserted into the T-DNA region of a binary vector such as pBIN19 and transformed into Arabidopsis. Agrobacterium-mediated transformation into Arabidopsis is performed by the vacuum infiltration or root transformation procedures known to the skilled man. Transformed seeds are selected and harvested and (where possible) transformed lines are established by normal selfing. Parallel transformations with 35S promoter-SERK constructs and the entire SERK gene itself are used as controls to evaluate over-expression in many cells or only in the few cells that naturally express the SERK gene. The 35S promoter-SERK construct may give embryo formation wherever the signal that activates the SERK-mediated transduction chain is present in the plant. A testing system based on emasculation and the generation of donor plant lines for pollen carrying LTP1 promoter-GUS and SERK promoter-barnase is established.

The same constructs (35S, EP3-1, AtChitlV, AtLTP-1 and SERK promoters fused to the SERK coding sequence) are employed for transformation into several *Arabidopsis* backgrounds. These backgrounds are wild type, male sterile, fis (allelic to emb 173) and primordia timing (pt)-1 lines, or a combination of two or several of these backgrounds. The wt lines are used as a control to evaluate possible effects on normal zygotic embryogenesis, and to score for seed set without fertilization after emasculation. The ms lines are used to score directly for seed set without fertilization. The fis lines exhibit a certain degree of seed and embryo development without fertilization, so may be expected to have a natural tendency for apomictic embryogenesis, which may be enhanced by the presence of the SERK constructs. The pt-1 line has superior regenerative capabilities and has been used to initiate the first stably embryogenic *Arabidopsis* cell suspension cultures. Combinations of several of the above backgrounds are obtained by

crossing with each other and with lines containing ectopic SERK expressing constructs. Except for the ms lines, propagation can proceed by normal selfing, and analysis of apomictic traits following emasculation. A similar strategy is followed in which the ATChilV, AtLTP-1 and SERK promoters are replaced by the bel-1 and fbp-7 promoters as well by other promoters specific for components of the female gametophyte.

Additional constructs are generated that have constitutive receptor kinase activity. Most of the receptor kinases of the SERK type act as homodimeric receptors, requiring autophosphorylation before being able to activate downstream signal transduction cascades. In many receptor kinases the extracellular domain serves as an inhibitor of the kinase domain in the ligand-free stage. This arrest is removed after binding of the ligand (Cadena and Gill, 1992). By introduction of a SERK construct, from which the extracellular ligand-binding domain has been removed, mutant homodimeric (in cells that do not have a natural population of SERK proteins) or neterodimeric (in cells that also express the unmodified forms) proteins can be generated with a constitutively activated kinase domain. This approach, when coupled to one of the promoters active in the nucellar region, results in activation of the embryogenic pathway in the absence of the activating signal. This may be an important alternative in cases where it is necessary or desirable to have activation of the SERK pathway only dependant on specific promoter activity and independent of temporal regulation of an activating signal. Introduction of SERK constructs that result in fertilization-independent-embryogenesis (fie) are tested in other species for their effect. In order to recognize the fie phenotype, the skilled man will use appropriate male sterile backgrounds. However, pollination is often necessary for apomixis of the adventitious embryony type, in order to ensure the production of endosperm.

Whilst the present invention has been particularly described by way of the production of apomictic seed by heterologous expression of the SERK gene in the nucellar region of the carpel, the skilled man will recognize that other genes, the products of which have a similar structure/function to the SERK gene product, may likewise be expressed with similar results. Moreover, although the example illustrates apomictic seed production in *Arabidopsis*, the invention is, of course, not limited to the expression of apomictic seed-inducing genes solely in this plant. Moreover, the present disclosure also includes the possibility of expressing the SERK (or related) gene sequences in the transformed plant material in a constitutive - tissue non-specific manner (for example under transcriptional control of a CaMV35S or NOS promoter). In this case, tissue specificity is assured by the localized presence within the vicinity of the embryo

sac of the ligand of the product of the said gene. Furthermore, the SERK (or related) gene products may interact with proteins such as transcription factors which are involved in regulating embryogenesis. This interaction within tissue which has been transformed according to the present disclosure is also part of the present invention.

PCT/EP97/02443

The skilled man who has the benefit of the present disclosure will also recognize that the SERK gene (and others as indicated in the preceding paragraph) may be transformed into plant material which may be propagated and/or differentiated and used as an explant from which somatic embryos can be obtained. Expression of such sequences in the transformed tissue (which is subjected to a ligand of the kinase gene products) substantially increases the percentage of the cells in the tissue which are competent to form somatic embryos, in comparison with the number present in non-transformed like tissue.

The invention will be further apparent from the following description and the associated drawings and sequence listings.

SEQ ID NO. 1 depicts the *Daucus carota* genomic clone of the putative receptor kinase (SERK) associated with the transition of competent to embryogenic cells;

SEQ ID NO. 2 depicts the cDNA of the said putative kinase;

SEQ ID NOs. 3 depicts the the predicted protein sequence of the SERK protein encoded by the DNA of SEQ ID NO:1.

SEQ ID NOs: 4-16 depict the sequences of various PCR primers; and

SEQ ID NOs. 17-19 depict specific peptides contained within the gene product of SEQ ID NO. 2.

SEQ ID NO: 20 depitcts the *Arabidopsis thaliana* partial genomic clone of the putative receptor kinase (SERK) associated with the transition of competent to embryogenic cells.

SEQ ID NO: 21 depicts the predicted protein sequence of the SERK protein encoded by the DNA of SEQ ID NO:20.

SEQ ID NOs: 22, 24, 26, 28 and 30 depict the partial DNA sequences of 5 EST clones with high homology to the SERK LRR sequences.

SEQ ID NOs. 23, 25, 27, 29 and 31 depict the predicted protein sequence of the partial DNA sequences of the 5 EST clones of SEQ ID Nos: 22, 24, 26, 28 and 30.

SEQ ID NO: 32 depicts the nuclotide sequence of the SERK cDNA from Arabidopsis thaliana.

SEQ ID NO: 33 depicts the predicted amino acid sequence of the SERK protein from Arabidopsis thaliana encoded by the DNA of SEQ ID NO: 32. Figure 1 shows the results of an RT-PCR experiment performed on RNA extracted from the indicated tissues. 40 cycl s followed by Southern blotting of the resulting bands is necessary to visualize SERK expression. Lanes include explants at day 7, treated for less (lane 1) or more (lane 2) then 3 days with 2,4-D. In the original a very faint signal is visible in lane 2, but not in lane 1. Established embryogenic cultures (lanes 4-6) but not a non-embryogenic control (lane 3) express the SERK gene. In carrot plants, no expression is detectable except for developing seeds after pollination (lane 7). Up to day 7 after pollination, the carrot zygote remains undivided, suggesting that the observed signal is coming only from the zygote. At day 10, the early globular and at day 20 the heart stage is reached in carrot zygotic embryogenesis. No signals are seen on Northern blots.

Figure 2A shows the results of a whole-mount *in situ* hybridization with the SERK cDNA on 7 day explants treated for 3 days with 2,4 D. Few cells on the surface of the explant express the SERK gene, and those cells that do are the ones that become embryogenic. Figure 2B shows a whole mount *in situ* hybridization on a partially dissected seed containing a globular zygotic embryo. Hybridization is visualized by DIG staining.

Figure 3 shows SERK expression in embryogenic hypocotyl cells during hormone-induced activation, determined by whole mount *in situ* hybridization. Bar: 50 mm

- (A-E) Cell population generated by mechanical fragmentation of the activated hypocotyls. Only few of a certain type of cell, defined enlarged cell show SERK expression (asterisks). Small cytoplasmic cells (c), enlarging cells (eg) and large cells (l) never show SERK expression.
- (F) Hypocotyl longitudinal section before hormone-induced activation. It is not possible to detect any SERK expression in any type of cell.
- (G-I) Proliferating mass coming from the inner hypocotyl tissues 10 days after the beginning of the hormonal treatment (longitudinal section). In G a single enlarged cells showing SERK expression is detectable within a row of negative cells showing the same morphology. In H a single enlarged cell showing serk expression is detaching from the surface of the proliferating mass. In I a cluster of enlarged cells showing SERK expression is detectable at the surface of proliferating tissue.
- (J) Proliferating mass coming from the inner tissues of the hypocotyl 10 days after the beginning of the rooting treatment (24 hours with 2,4-D followed by homone removal). Both the root primordia and the enlarged cells detaching from the surface do not show any SERK expression.

WO 97/43427 PCT/EP97/02443

Figure 4 shows the phenotype of Arabidopsis WS plants transformed with the 2200 bp SERK-luciferase consturct at the seedling level. Pictures were taken at 28 days after germination of T2 seeds. In plant II and III no clear shoot meristem is visible at the seedling stage, 7 days after germination. The first two leaves, if they develop at all, are needleshaped as hown on the pictures taken 28 days after germination. At this time plant I, which shows no clear phenotype, already starts flowering. Secondary shoot meristems are already developing in plant no II and will also develop later from no III. Shoot meristems, influorescences and normal flowers eventually develop on all plants.

Figure 5 shows how the 2200 bp SERK luciferase construct affects the number of developing ovules in the siliques of transformed plants.

Figure 6 shows autophosphorylation of purified SERK fusion protein *in vitro*. Lane 1: purified SERK fusion protein; Lane 2: serine phosphate; Lane 3: threonine phosphate; Lane 4: thyrosine phosphate.

The following description illustrates the isolation and cloning of the SERK gene and the production of apomictic seed by heterologous expression of the said gene in the nucellar region of the carpel so that somatic embryos form which penetrate the embryo sac and are encapsulated by the seed as it develops.

ISOLATION AND CLONING OF THE SERK GENE FROM DAUCUS CAROTA

Isolation of cDNA clones that are preferentially expressed in embryogenic cell cultures of carrot

In order to increase the chance of success for obtaining genes expressed in carrot suspension cells competent to form embryos, the number of embryo-forming cells as present in a series of established cell cultures was determined. A sub-population of cells that passed through a 30 mm nylon sieve was isolated from eight different cultures that ranged in age between 2 months and 4 years. In these sub 30 mm populations, the number of embryos formed from the single cells and small cell clusters was determined and expressed as a percentage of the total number of cells present at the start of embryogenesis. Sieved <30 mm cultures able to form somatic embryos with a frequency of more than 1% were then used as a source for competent cells, and cultures that produced less than 0.01% embryos were used as non-embryogenic controls. As main cloning strategies, cold plaque screening (Hodge et al. 1992) and differential display (dd) RT-PCR (Liang and Pardee, 1992) were used besides conventional differential screening of cDNA libraries.

Labeled probes for differential screening were obtained from RNA out of a <30 mm sieved sub-population of cells from either embryogenic or non-embryogenic cell cultures. Employing these probes in a library screen of approximately 2000 plaques yielded 26 plaques that failed to show any hybridization to either probe. These so-called cold plaques were purified and used for further analysis. From the total number of plagues that did hybridize, about 30 did so only with the probe from embryogenic cells, ddRT-PCR reactions using a combination of one anchor primer and one decamer primer were performed on mRNA isolated from three embryogenic, and three non-embryogenic suspension cultures. About 50 different ddRT-PCR fragments were obtained from each reaction. Using combinations of three different anchor and six different decamer primers, a total of approximately 1000 different cDNA fragments was visualized. Six of these PCR fragments were only found in lanes made with mRNA from <30 mm populations of cells from embryogenic cultures (Table 1) and with oligo combinations of the anchor primer (5'-TTTTTTTTTGC-3') and the decamer primers (5'-GGGATCTAAG-3'), (5'-ACACGTGGTC-3'), (5'-TCAGCACAGG-3'). Because differential PCR fragments often consist of several unresolved cDNA fragments (Li et al. 1994), cloning proved to be essential prior to undertaking further characterization of the PCR fragments obtained.

All clones obtained were subjected to a second screen, that consisted of spot-dot Northern hybridization performed under conditions of high stringency. This method, that used RNA from entire unsieved embryogenic and non-embryogenic suspension cultures, proved to be a fast and reliable additional selection method. Only one clone (22-28) of the 30 clones obtained after differential screening, proved to be restricted to embryogenic cell cultures while the majority was constitutively expressed. The 26 clones obtained from the cold plaque screening required long exposure times in the spot-dot Northern analysis. Six of these clones failed to show any hybridization signal and 19 proved to be expressed in both embryogenic and non-embryogenic cell cultures. One clone (31-50) showed low expression in all embryogenic cultures, and in one non-embryogenic culture, but not in the others. Of the six cloned fragments obtained by ddRT-PCR display, four showed hybridization more or less restricted to transcripts present in embryogenic cultures. All clones that passed through the second screening were sequenced. Two of the ddRT-PCR clones (6-8 and 7-13) were identical to the carrot Lipid Transfer Protein (LTP) gene, previously identified as a marker for embryogenic carrot cell cultures. LTP expression is restricted to embryogenic cell clusters and the protoderm of somatic and zygotic embryos from the early globular stage onwards (Sterk et al. 1991). Therefore, while the LTP gene is not a marker for competent cells, its appearance in the screening confirms the validity of our methods with respect to the cloning of genes expressed early during somatic embryogenesis.

cDNA clone 31-50 encodes a leucine-rich repeat containing receptor-like kinase

The mRNA corresponding to the isolated clone 31-50 had an open reading frame of 1659 nucleotides encoding a protein with a calculated Mw of 55 kDa. Because clone 31-50 is mainly expressed in embryogenic cell cultures it was renamed Somatic Embryogenesis Receptor Kinase (SERK). The SERK protein contains a N-terminal domain with a five-times repeated leucine-rich motif that is proposed to act as a protein-binding region in LRR receptor kinases (Kobe and Deisenhofer, 1994). Between the extracellular LRR domain of SERK and the membrane-spanning region is a 33 amino acid region rich in prolines (13), that is unique for th SERK protein. Of particular interest is the sequence SPPPP, that is conserved in extensins, a class of universal plant cell wall proteins (Vamer and Lin, 1989). The proposed intracellular domain of the protein contains the 11 subdomains characteristic of the catalytic core of protein kinases. The core sequences HRDVKAAN and GTLGYIAPE in respectively the kinase subdomains VB and VIII suggest a function as a serine / threonine kinase (Hanks *et al.* 1988).

Another interesting feature of the intracellular part of the SERK protein is that the C-terminal 24 amino-acids resembles a single LRR. The serine and threonine residues present within the intracellular LRR sequence are surrounded by acidic residues and might be targets for the autophosphorylation of SERK, thereby regulating the ability of other proteins to interact with this receptor-kinase in a similar fashion as described for the SH2 domain of the EGF family of tyrosine receptor kinases.

Hybridization of the SERK cDNA clone to the carrot genome revealed the presence of only a single main hybridizing band after digestion with EcoR1, probably reflecting a single SERK gene in the carrot genome. This was confirmed after digestion with Ddel, an enzyme that cuts three times within the SERK gene. No signal was observed after Northern blotting of mRNA from embryogenic cell cultures and hybridization with labeled SERK probes, reflecting the low levels of transcript present in these cultures. Detection of the SERK transcript on the original spot-dot Northerns was only possible after long exposure times compared with other probes.

The ability of the SERK protein to autophosphorylate was investigated *in vitro*, using a previously described autophosphorylation assay (Mu et al. 1994), with a bacterial fusion protein that contained the complete intracellular region of the SERK protein. The bacterially expressed SERK fusion protein was able to autophosphorylate, indicating that the SERK protein is able to fulfill a role as a protein kinase *in vivo* (Heldin, 1995).

Expression of the SERK gene corresponds with the first appearance of competent cells during hypocotyl activation

When carrot hypocotyls are induced with 2,4-D, only the cells of the provascular tissue proliferate. Cells of epidermal and cortical origin merely expand, suggesting that the provascular tissue derived cells form the newly initiated suspension culture. After removal of 2,4-D, the formation of somatic embryos occurs after 2-3 weeks. Somatic embryos are preceded by embryogenic cells, that are developed in turn from competent cells. While competent and embryogenic cell formation take place in the presence of 2,4-D, it was not clear when this occurred, and which cells acquired competence. Since previous experiments (Toonen et al. 1994) revealed that cell morphology is not a good criterion, the first appearance of single competent cells was determined experimentally by semi-automatic cell tracking performed on large populations of immobilized cells. Hypocotyl explants activated with 2,4-D for seven days were mechanically fragmented and samples of the r sulting population of mainly single

suspension cells were immobilized to allow recording of their development by cell tracking. In the immobilized cell populations obtained in this way all the morphologically discernible cell types were present that were also seen in the un-fragmented activated hypocotyls. Because the different cell morphologies observed during hypocotyl activation were known (Guzzo et al. 1995), it was possible to trace back the original position of each type of cell in the activated explant. Small cytoplasm-rich cells (16x16 mm) are the proliferating cells that surround the vascular elements. Enlarging vacuolated cells (16x40 mm) are encountered on the surface of the mass of proliferating cells and these can detach from the surface when fully enlarged (35x90 mm). Large vacuolated cells (more than 60x140 mm) are the non-proliferating remnants of the hypocotyl epidermis and cortical parenchyma. The shape of the enlarging and fully enlarged cells could change from oval to elongate or triangular. Cell tracking on a total of 24,722 cells released from seven days activated hypocotyls showed that only 20 single cells formed a somatic embryo. Because of their dependance on continued 2,4-D treatment, the embryoforming single cells are still in the competent cell stage. All of the embryo-forming single cells belonged to the category of 3.511 enlarged cells, that contained therefore competent cells in a frequency of 0.56%. The single cell tracking experiments clearly reveal that the ability of explant cells to reinitiate cell division under the influence of 2,4-D, resulting in a population of highly cytoplasmic and rapidly proliferating cells, does have a causal relation with the ability to become embryogenic. It is also clear that only a very limited number of the cells that make up the newly initiated embryogenic suspension culture are actually competent to form embryogenic cells.

Expression of the SERK gene, determined by whole mount *in situ* hybridization on a similar population of cells as used for the cell tracking experiments, was found to be restricted to only 0.44% of the enlarged cells. Therefore, the expression of the SERK gene appears closely correlated both qualitatively and quantitatively with the presence of competent single cells.

To obtain insight into the temporal regulation of SERK expression in the course of explant activation, whole mount *in situ* hybridization was performed on entire intact or hand-sectioned explants treated for different periods with 2,4-D. Representative samples were collected at daily intervals from explants untreated and treated for three days, six days, seven days or ten days with 2,4-D before returning to B5-0. No SERK-expressing cells were ever found in explants treated for less then three days with 2,4-D. While enlarged cells became present after the first five days of culture, the first few SERK-expressing enlarged cells were found after six-seven days of culture in the presence of 2,4-D treatment. These few cells were present at the surface

of the mass of proliferating cells originating from the provascular tissue. In the hypocotyls treated for ten days with 2,4-D, the number of SERK-positive cells had increased to 3.04% and included at this stage also cells present in small clusters. No SERK transcript was ever detected in small cytoplasm-rich cells or large vacuolated cells. Hypocotyls were also treated for only one day with 2,4-D and subsequently cultured in hormone-free medium for a total of seven or ten days. Under these conditions explant cells proliferated and gave rise to roots and non-embryogenic cell cultures, while SERK expression could never be detected. The in situ hybridization results described above were obtained from a relatively small number of explants and a few hundred cells, so RT-PCR followed by Southern hybridization was performed to obtain more quantitative results. These are shown in Figure 7 and confirm the close temporal correlation between the first appearance of competent cells in explants treated for three days with 2,4-D and the expression of the SERK gene. Northern hybridization never gave any signal after hybridization with SERK cDNA probes, not even after prolonged exposure in a Phosphorlmager, in line with the extremely restricted expression pattern of the SERK gene.

Expression of the SERK gene corresponds with the occurrence of competent cells in established embryogenic cell cultures

While the results described so far indicate that competent and embryogenic cell formation is restricted to a particular class of enlarged cells during explant activation, the situation in an established embryogenic cell culture is more complex. Competent single cells in such cultures do not appear to belong to one cell type in particular, but have been shown to originate from all morphologically different cell types. In cell tracking experiments, embryogenic cells, that do not require exogenous auxin treatment, were never observed to be single but consisted of clusters of at least 3-4 cells (Toonen et al. 1994). SERK expression was found in all morphologically discernible single cell types that were present in an embryogenic cell culture at a frequency between 0.1 and 0.5% depending on the cell type. In non-embryogenic cultures, SERK expression was not restricted to single cells, but also occurred in small clusters of 2 to 16 cells. Since clusters of this size are known to consist of embryogenic cells, these data show that SERK expression is not restricted to competent single cells, but may persist in small clusters of embryogenic cells. No SERK expression was encountered during the late globular, heart and torpedo-stages of somatic embryogenesis.

Th SERK g ne is transi ntly express d in zygotic embryogenesis

Th expression of the SERK gene in carrot plants was determined by RT-PCR. The results indicate that no SERK mRNA accumulates in any of the adult plant organs nor in flowers prior to pollination. The first occasion when SERK expression can be detected is in flowers at three days after pollination (DAP), at which stage fertilization has taken place and endosperm development has commenced. SERK mRNA remains present in flowers up to twenty DAP, corresponding with the early globular stage of the zygotic embryo (Yeung et al. 1996). Whole mount in situ hybridization on partially dissected carrot seeds confirmed that the SERK gene was only expressed in early embryos up to the globular stage. Expression was observed in the entire embryo including the suspensor. No expression was seen in seedlings, roots, stems, leaves, developing and mature flower organs, pollen grains and stigma's before and after fertilization. Tissues in the developing seed such as seed coat, integuments, all embryo sac constituents before fertilization as well as the endosperm at all stages of development investigated did not show any SERK expression. Later stages of carrot zygotic embryos were also completely devoid of SERK mRNA. Given this pattern of expression, that is restricted to the zygotic embryo, the signal as detected by RT-PCR in flowers at 3 and 7 DAP must come from SERK mRNA as present in zygotes, because in carrot the zygote remains undivided up to one week after pollination (Yeung et al. 1996). Although SERK expression persists to slightly later stages in zygotic globular embryos when compared to the somatic ones, these results confirm the transient pattern of expression as observed for the SERK gene during somatic embryogenesis and also imply that there is a correspondence between the formation of competent cells in vitro and the formation of the zygote in vivo.

METHODS

Cell culture, hypocotyl explant induction and cell tracking

Cell cultures were derived from *Daucus carota* cv. Flakkese and maintained as previously described (De Vries *et al.* 1988a). Cell suspension cultures were maintained at high cell density in B5 medium (Gamborg *et al.* 1968) supplemented with 2 mM 2,4-D (B5-2 medium). Embryo cultures with globular, heart and torpedo-stage somatic embryos were derived from <30 mm sieved cell cultures cultured at low cell density (100 000 cells / ml) in B5 medium without 2,4-D (B5-0). For hypocotyl explant induction experiments, plantlets were obtained from seed of

Daucus carota cv. S Valery as described previously (Guzzo et al., 1994). The hypocotyls of one w ek old plantlets were divided in segments of 3-5 mm, incubated for various periods of time in B5-2 medium and returned to B5-0 medium. Seven days after explantation and exposure to 2,4-D the hypocotyl segments were fragmented on a 170 mm sieve and the resulting cells collected to form a fine cell suspension. Immobilization of these cells in B5-0.2 medium was performed in a thin layer of phytagel (Toonen et al. 1994). After one week of further culture 2,4-D was removed by washing the plates with B5-0 medium. This allowed embryos to develop beyond the globular stage. Recording the development of the immobilized cells was performed with a procedure modified from the previously described by Toonen et al. (1994). The main change involved a new MicroScan program for automatic 3-axis movement to scan all cells in the phytagel (Toonen et al. 1996).

Nucleic acid isolation and analysis

RNA was isolated from cultured cells and plant tissues as described by De Vries *et al.* (1988b). Poly(A)*-RNA was obtained by purification by oligo (dT) cellulose (Biolabs). For RNA gel blot analysis samples of 10 mg total RNA were electrophoresed on formamide gel, and transferred to nytran-plus membranes. For RNA spot-blot analysis 5 mg of total RNA was denatured and spotted onto nytran-plus filters using a hybridot manifold (BRL).

Genomic DNA was isolated according to Sterk et al. (1991). Samples of 10 mg genomic DNA were digested with different restriction enzymes and separated on agarose gel, and transferred to nytran-plus membrane (Schleicher & Schuell). Hybridization of RNA blots took place at 42°C in hybridization buffer containing 50% formamide, 6xSSC, 5xDenhardt, 0.5% SDS and 0.1 mg/ml salm sperm DNA. Hybridization of DNA blots was performed as previously described (Sterk et al. 1991). Following hybridization, filters were washed under stringent conditions (3x20 min in 0.1% SSC, 1% SDS, at 65°C). Filters were exposed to Kodak X-Omat AR film. The integrity and the amount of RNA on the blots was confirmed by hybridization with an 18S ribosomal RNA probe. Nucleotide sequence analysis was performed on an ABI 373A automated DNA sequencer (Applied Biosystem).

Screening procedures

Two independent cDNA libraries were constructed with equal amounts of poly(A)*-RNA from total established cell cultures grown for six days in B5-2 medium, sieved <125 mm cell cultures grown for six days in B5-0 medium and sieved <30 mm cell cultures grown for six days in B5-0

WO 97/43427 PCT/EP97/02443
- 25 -

medium. cDNA synthesis and cloning into the Uni-ZAPTM XR vector was performed according to the manufacturers protocol (Stratagene).

Differential screening of the cDNA libraries was performed essentially as described by Scott et al. (1991). RNA was isolated from either three embryogenic or three non-embryogenic cell cultures, that were grown for seven days in B5-2 after sieving through 30 mm mesh. First strand cDNA synthesis was performed on 4 mg total RNA using AMV reverse transcriptase (Gibco BRL). [³²P]dATP labeled probes were prepared using random prime labeling on first strand cDNA. Pooled probes from embryogenic and non-embryogenic cell populations were hybridized to two pairs of nitrocellulose filters, each containing 1000 plaques from one cDNA library. After washing for 3x20 min in 0.1% SSC, 1% SDS at 65°C, hybridization was visualized by autoradiography for two days on Kodak X-omatic film. Plaques that only showed signal with the embryogenic transcript probe were purified by two further rounds of screening.

In order to identify cDNA clones which are expressed at low levels in the <30 mm sieved cell population, cold plaque screening was performed as described by Hodge *et al.* (1992). Plaques from the differential screening that did not show any signal after seven days of autoradiography were purified by two further rounds of screening. The resulting clones were used as probes for characterization of the expression pattern of the corresponding genes.

Differential Display RT-PCR

Differential display of mRNA was performed essentially as described by Liang and Pardee (1992). cDNA synthesis took place by annealing 1 mg of total RNA in 10 ml buffer containing 200 mM KCl, 10 mM Tris-HCl (pH 8.3), and 1 mM EDTA with 100 ng of one of the following anchor primers: (5'-TTTTTTTTTTTGC-3'), (5'-TTTTTTTTTCTG-3'), (5'-TTTTTTTTTCA-3'). Annealing took place by heating the mix for 3 min. at 83°C followed by incubation for 30 min at 42°C. Annealing was followed by the addition of 15 ml pre-warmed cDNA buffer containing 16 mM MgCl₂, 24 mM Tris-HCl (pH 8.3), 8 mM DTT, 400 mM dNTP, and 4 Units AMV reverse transcriptase (Gibco BRL). cDNA synthesis took place at 42°C for 90 min. First strand cDNA was phenol/chlorophorm extracted and precipitated with ethanol using glycogen as a carrier. The PCR reaction was performed in a reaction volume of 20 ml containing 10% of the synthesized cDNA, 100 ng of anchor primer, 20 ng of one of the following 10-mer primers: (5'-GGGATCTAAG-3'), (5'-TCAGCACAGG-3'), (5'-GACATCGTCC-3'), (5'-CCCTACTGGT-3'), (5'-GCGTGGTC-3'), (5'-GGTGACTGTC-3'), 2 mM dNTP, 0.5 UnitTag enzyme in PCR buffer (10

mM Tris-HCI (pH 9.0), 1.5 mM MgCl₂, 50 mM KCI, 0.01% gelatin and 0.1% Triton X100) and 6 nM [a-32P] dATP (Amersham). PCR parameters were 94°C for 30 sec, 40°C for 1 min, and 72°C for 30 sec for 40 cycles using a Cetus 9600 (Perkin-Elmer). Amplified and labeled cDNAs were separated on a 6% denaturing DNA sequencing gel. Gels were dried without fixation and bands were visualized by 16 hours of autoradiography using Kodak X-omatic film. Bands containing differentially expressed cDNA fragments of 150-450 nucleotides were cut out of the gel and DNA was extracted from the gel slices by electroelution onto DE-81 paper (Whatmann). After washing of the paper in low salt buffer (100 mM LiCl₂ in 10 mM TE buffer), and elution of the cDNA in high salt buffer (1 M LiCl2 in 10 mM TE buffer with 20% ethanol) the cDNA was concentrated by precipitation in ethanol using glycogen as carrier. Reamplification of the cDNA fragments using the same PCR cycling parameters as described above but PCR buffer containing 2.5 mM of both the 10-mer and the anchor oligo and 100 mM dNTP, DE-81 paper allowed an efficient recovery of the DNA fragments and reamplification generated an average of 500 ng DNA after 40 cycles. Amplified PCR products were blunt-ended using the Klenow fragment of E.coli DNA Polymerase I (Pharmacia), purified on Sephacryl-S200 columns (Pharmacia), ligated into a Smal linearized pBluescript vector II SK (Stratagene) and transformed into E.coli using electroporation.

RT-PCR

Adult plant tissues from *Daucus carota* were obtained from S&G Seeds (Enkhuizen). Controlled pollination was performed by hand. Flower tissue RNA was obtained from three compete umbels for each time-point and contained all flower organs including pollen grains. 2 mg of total RNA from adult plant tissue or cell cultures was annealed at 42°C with 50 ng oligo (5'-TCTTGGACCAGATAATTC-3') in 10 ml annealing buffer (250 mM KCl, 10 mM Tris-HCl pH 8.3, 1 mM EDTA). After 30 min. annealing, 1 unit AMV-reverse transcriptase was added in a volume of 15 ml cDNA buffer (24 mM Tris-HCl pH 8.3, 16 mM MgCl₂, 8 mM DTT, 0.4 mM dNTP). The reverse transcription reaction took place for 90 min. at 42°C. PCR amplification of SERK-cDNA was carried out with two specific oligos for the SERK kinase domain, (5'-CTCTGATGACTTTCCAGTC-3') and (5'-AATGGCATTTGCATGG-3'). Amplification was carried out with 30 cycles of 30 sec. at 94°C, annealing at 54°C for 30 sec. and extension at 72°C for 1 min., followed by a final extension for 10 min.at 72°C.

Wh le mount in situ hybridization

Whole mount *in situ* hybridizations were performed essentially as previously described (Engler et al. 1994). Cell cultures and somatic embryos were immobilized on poly-L-lysine coated glasses during fixation to improve handling. Whole mount *in situ* hybridization on explants took place by embedding hypocotyls from seven-days old plantlets in 3% Seaplaque agarose (Duchefa) and processing them in Eppendorf tubes. Transverse as well as longitudinal sections were made with a vibrotome (Biorad Microcut). Sections of 50-170 mm thick were incubated in B5-2 medium for a minimum of three days to induce formation of embryo-forming cells. Optimal induction was achieved with longitudinal hypocotyl sections with a thickness of at least 90 mm. To obtain proliferating, non-embryogenic cell cultures, hypocotyl sections were exposed to 2,4-D for only 1 day, and subsequently transferred to B5-0 medium (Guzzo *et al.* 1994). Whole mount in situ hybridization on developing seeds was performed by removing the chalazal end of the seeds to allow easier probe penetration. After hybridization, the enveloping layers of integuments and endosperm were carefully removed to expose the developing embryos. In situ hybridization on sections was performed as described previously (Sterk et al. 1991) except for the use of non-radioactive probes.

All samples were fixed for 60 min. in PBS containing 70 mM EGTA, 4% paraformaldehyde, 0.25% glutaraldehyde, 0.1% Tween 20, and 10% DMSO. Samples were then washed, treated with proteinase K for 10 min, again washed and fixed a second time. Hybridization solution consisted of PBS containing 0.1% Tween 20, 330 mM NaCl, 50 mg/ml heparin, and 50% deionized formamide. Hybridization took place for 16 hours at 42°C using digoxigenin-labeled sense or antisense riboprobes (Boehringer Mannheim). After washing the cells were treated with RNaseA, and incubated with anti-digoxigenin-alkaline phosphatase conjugate (Boehringer Mannheim) which had been preabsorbed with a plant protein extract. Excess antibody was removed by washing followed by rinsing in staining buffer (100 mM Tris-HCl pH 9.5, 100 mM NaCl, 5 mM MgCl₂, 1 mM levamisole) and the staining reaction was performed for 16 hours in a buffer containing NBT and BClP. Observations were performed using a Nikon Optiphot microscope equipped with Nomarski optics.

Autophosphorylation assay

A 1.4 kB Sspl cDNA fragment of the SERK cDNA encoding most of the open reading frame apart from the N-terminal three LRRs was cloned into the pGEX expression vector (Pharmacia).

A fusion protein consisting of SERK and the glutathione S-transferase gene product was synthesized by a three hours induction of transformed *E.coli* with 2 mM IPTG. Fusion protein was isolated and purified as described previously (Hom and Walker, 1994). Purified fusion protein was coupled to glutathione agarose beads (Sigma) and incubated for 20 min. at 20°C in a volume of 10 ml buffer: 50 mM Hepes (pH 7.6), 10 mM MgCl₂, 10 mM MnCl₂, 1 mM DTT, 1 mCi [y -³²P] (3 000 Ci/mmol) . Excess label was removed by washing the fusion protein/glutatione agarose beads three times for 5 min. in 50 mM Tris-HCl (pH 7.3), 10 mM MgCl₂ at 4°C. Protein was removed from the beads by cooking in SDS-PAGE loading buffer. Equal amounts of protein were separated by SDS-PAGE and protein autophosphorylation was visualized by autoradiography.

SERK fusion proteins produced with the Baculovirus expression system.

Further fusion proteins containing the intracellular part of the *Daucus carota* SERK protein (1.0 kB HindIII / Sspl fragment of the carrot SERK cDNA clone 31-50) were made using the baculorvirus vector pAcHLT.

In vitro phosphorylation studies with this purified protein showed that most if not all of the autophosphorylation of this SERK fusion protein was at threonine residues (Figure 6)

Construction of viral transfer vectors

The pAcHLT-B and pAcHLT-C baculovirus transfer vectors were used for the cloning of two cDNA fragments of the carrot SERK gene. The Sspl 1.41 kB fragment of carrot DcSERK cDNA was cloned into the Smal site of pAcHLT-B and the Sspl / Pvull 1.07 kB fragment of carrot DcSERK cDNA was cloned into the Smal site of pAcHLT-C. The first construct contains the complete C-terminal part of the DcSERK protein and from the putative extracellular region the proline-rich region and three of the lecuine-rich repeats. The second construct contains only the putative intracellular region of the DcSERK gene product. Nucleotide sequence analysis was performed in order to confirm the presence and the orientation of the DcSERK cDNA within the vector.

Transformation of insect cells

The resulting transfer vectors were used to transfect (lipofect) insect cell culture Sf21 from Spodoptera frugiperda in combination with linearized AcMNPV baculovirus DNA. Monolayers of SF21 cells were transfected in 35 mm petridishes containing 2 ml of Hink's

WO 97/43427 PCT/EP97/02443 - 29 -

medium. One microgram of linearized AcMNPV baculovirus DNA (Baculogold, Invitrogen) was added to 5 microgram of pAcHLT / SERK vector construct in 25 microliter of water. Fifteen microliter of Lipofectin (BRL) was mixed with 10 microliter of water, after which the DNA solution was added. After mixing 200 microliter of Hink's medium was added to the mix and the solution was transferred to the cell monolayer, from which the medium was removed. After one hour, 500 microliter of Hink's medium was added and the cells were incubated for anotehr 3 hours. Finally, 1 ml of Hink's medium with 20% foetal bovine serum (FBS) was added and the cells were incubated for 4 days. After transfection, the viral infection could be identified by the reduced growth of cells, the swollen shape and the enlarged nucleus. After four days, infected cells were harvested and the medium containing infectious budded virus was collected and used for plaque assays and amplification of recombinant virus stocks.

Isolation of single recombinant viruses

Single recombinant virus plaques were isolated from monolayers of cells infected with a titration range of the primairy virus stock. Infections was performed in 35 mm petridishes with monolayers of cells. Virus stocks were diluted in 600 microlieter of Graces medium and added to the cell monolayer, followed by a 90 minutes incubation period at in Graces medium with 20% FBS. Afterwards, 3% Sea Plaque agarose was autoclaved, mixed with an equal amount of 2x Graces medium with 20% FBS and from the resulting agarose overlay solution 2 ml. was spread over the cell monolayers after removal of the viral inoculum. After 4 days of incubation single plaques could be visualized and purified for further analysis.

Fusion protein production.

After determining the titer of purified recombinant viruses, monolayers of Sf21 cells in 75 cm² flasks were infection with a multiplicity of infection (MOI) of 10. Incubation of cells with the virus inoculum was performed for 90 min. after which 8 ml. of Hink's medium with 10%FBS was added. After 3 days of incubation, cells were harvested and washed twice with PBS. Cells were lysed for 45 min on ice in twenty volumes of 1x insect cell lysis buffer (10 mM Tris pH 7.5, 130 mM NaCl, 1% Triton, 100 mM NaF, 10 mM NaPi, 10 mM NaPPi, with proteinase inhibitors: 16 mg/l benzamidine, 10 mg/l phenanthroline, 10 mg/l aprotinin, 10 mg/l leupeptin, 10 mg/l pepstatin A, 1 mM PMSF).

The lysate was cleared by centrifugation at 10.000 g for 30 min and the supernatant was batchwis incubated in TALON resin (with high affinity for the 6xHIS tag of the recombinant fusion protein). Binding was performed by gentle agitation for 20 min. at room temp. The resin was washed three times with lysis buffer, followed by an elution step with lysis buffer with 200 mM imidazole. Purified fusion protein was collected and purifty and integrity was tested by SDS-PAGE.

Autophosphorylation assays

Protein kinase activity was determined by incubating 1 microgram of purified fusion protein for 30 min. at room temp. in a buffer containing 10 mM MgCl2, 10 mM MnCl2, 1 mM DTT and 10 µM [gamma-32]ATP (10⁵ pm/pmol ATP). The autophosphorylated fusion protein was purified after SDS-PAGE from the gel in a buffer containing 50 mM NH4CO3, 0.1% SDS, 0.25% beta-mercaptoethanol. Protein was precipitated with 20 µg/ml BSA and 20% (w/v) solid trichloroacetic acid. The precipitate was collected after centrifugation, hydrolysed in 50 µl 6N HCl for 1 hour at 120 degrees Celcius. HCl was subsequently removed by lyophilization and the pellet was resuspended in a buffer consiting of 2.2% formic acid and 7.8% acetic acid. Hydrolysed protein was loaded onto cellulose thin layer chromatography plates together with control amino acid samples (phosphoserine, phosphothreonine, phosphotyrosine). Chromatography was performed in a buffer containing propionic acid: 1M ammonium hydroxide: isopropyl alcohol (90:35:35 v/v/v). After separation and drying of the plates, the separated amino-acids were visualized by spraying with 0.25% ninhydrin in aceton, followed by heating for 5 min. at 65 degrees Celcius. Plates were afterwards exposed to Phospho Imager casettes in order to detect the phospho-labeled aminoacids.

SERK antibodies

Purified fusion proteins (10 µg) were mixed in complete Freund adjuvant and injected IP into BALBc mice. After 4 weeks booster antigen was injected (10 µg purified fusion protein in imcomplete Freund adjuvant). Two weeks later a final booster was injected. One week after the final booster, serum was collected from these mice. The specificity and the titer of the resulting sera was tested on Western blots using total insect cell extracts with or without the SERK fusion proteins.

INTRODUCTION OF THE SERK GENE INTO *PLANTA* AND THE PRODUCTION OF APOMICTIC SEED

Carrot transformation with a SERK promoter fragment/luciferase gene fusion

The binary vector pMT500 is based on the pBIN19 vector (Bevan, 1984) and contains the firefly luciferase gene downstream of a polylinker containing 5 unique restriction sites was created by uni-directional ligation of the firefly luciferase coding region followed by the polyadenylation sequence from the pea *rbcS*::E9 gene in the *HindIII-XbaI* site of the binary vector pMOG800 (kindly provided by Mogen N.V., Leiden, The Netherlands). The binary vector pMOG800 is based upon pBIN19 (Bevan, 1984) but while in pBIN19 the polylinker is flanked by the left border and the neomycin phosphotransferase (NPT II) expression cassette, the polylinker in pMOG800 is flanked by the right border and the NPT II expression cassette. From a genomic lambda clone, transcription regulating sequences from the carrot SERK gene were isolated by digestion with *HindIII* and DraI (SEQ ID No. 1), and cloned into the HindIII / SmaI sites of pBluescript SK+. From the resulting vector a KpnI / SstI fragment containing the SERK genomic DNA was isolated and cloned into the KpnI / SstI sites of the binary vector pMT500. The resulting DNA construct, pMT531, contained the 2200 bp genomic SERK DNA fragment as promoter sequence, the luciferase gene as vital reporter, and the E9 transcription terminator sequence.

The binary vector pMT531 was transformed by electroporation into Agrobacterium tumefaciences strains MOG101 and MOG301 (for transformation into carrot cells) and into Agrobacterium tumefaciences strain C58C1 (for transformation into Arabidipsis thaliana plants). Transformed colonies were selected on LB plates with 100mg/l kanamycin.

Transformation of carrot cells

The firefly luciferase coding sequence under control of the genomic carrot HindIII / Dral 2200 bp DNA fragment was introduced into carrot cells by *Agrobacterium tumefaciens* mediated transformation of hypocotyl segments. Transformation of *Daucus carota cv*. 'Amsterdamse bak' was performed by slicing one week old dark grown seedlings into segments of 10 to 20 mm. Segments were incubated for 20 minutes in a freshly prepared 10 fold diluted overnight culture of *Agrobacterium*.. The segments were dried and transferred to a modified Gamborgs B5 medium (P1 medium; S&G seeds, Enkhuizen, The Netherlands) supplemented with 2 μ M 2,4-D (P1-2) and solidified with agar (Difco, Detroit, Mi, USA). After two days of culture in the dark at 25 \pm 0.5 _C, segments were transferred to

solidified P1-2 medium supplemented with kanamycin (100 mg·l⁻¹), carbenicillin (500 mg·l⁻¹; Duchefa) and vancomycin (100 mg·l⁻¹; Duchefa). After three weeks segments were transferred to fresh plates and transformed calli were selected after an additional three weeks. Transformed calli were grown on P1-2 plates with antibiotics for 3 weeks at a 16 hour light/8 hour darkness regime. Transformed embryogenic suspension cultures were initiated as described by transferring 0.2 g callus to 10 ml liquid P1-2 medium supplemented with 200 mg·l⁻¹ kanamycin, 250 mg·l⁻¹ carbenicillin and 50 mg·l⁻¹ vancomycin. During the first weeks 1 to 3 volumes of fresh medium were added to the culture at weekly intervals. After 5 to 7 weeks cultures were subcultured to a packed cell volume of 2 ml per 50 ml medium every two weeks and incubated at a 16 hour light / 8 hour darkness regime at 25 ± 0.5 °C.

- 32 -

One week after transfer to kanamycin selection medium, hypocotyl segments were sprayed with luciferin to test whether luciferase expression could be detected in transformed callus shortly after transformation. A large number of hypocotyl segments showed luciferase activity at the cut edges, but did not develop calli. Instead, growth of bacteria occurred, suggesting that the luciferase activity was of bacterial origin. Six to ten weeks after transformation, calli were obtained that showed luciferase activity in variable amounts, while no bacterial growth could be observed anymore. After 12 weeks, calli measuring 5 to 10 mm in diameter were used to start suspension cultures. At this time no bacterial contamination was observed. A control transformation experiment in which luciferase expression under influence of the CaMV 35S promoter was observed in single cells and cell clusters in the suspension culture demonstrating that the luciferase protein is active in Daucus carota suspension cultured cells.

Cell immobilisation

One-week old high-density (10⁶ - 10⁷ cells·ml⁻¹) suspension cultures were sieved through nylon sieves with successive 300, 125, 50 and 30 µm pore sizes (Monodur-PES; Verseidag Techfab, Walbeck, Germany). Single cells and cell clusters passing the last sieve are designated as < 30 µm populations. Control experiments with untransformed cells were performed with *Daucus carota* cv. 'Trophy' (S&G seeds) suspension cultures grown in P1-2 medium. Size fractionated cell populations smaller then 30 µm were immobilised in phytagel (P8196; Sigma, St Louis, Mo, USA) in petriperm dishes (Heraeus, Hanau, Germany). The bottom layer consisted of 1 ml P1-0 medium with 5 mM Ca ²⁺ and 0.2 % phytagel. Two

hundred thousand cells (< 30 μm and < 50 μm populations) in B5-0 medium without Ca ²⁺ supplemented with 0.1 % phytagel were poured on top of the bottom layer. For this layer B5 was applied since, at room temperature, phytagel solidified in P1 medium without Ca ²⁺. After 2 hours of solidification an additional P1-0 layer with 0.2 % phytagel was poured onto the cell layer preventing the B5 layer to move. To prevent dehydration of the phytagel layers and to supply luciterin to the cells, 0.5 ml P1-0 medium containing 0.05 μM luciferin (Promega, Madison, Wi, USA) was added after solidification. The final luciferin concentration in the culture was 0.02 μM. Luciferin detection on single cells was determined

Arabidopsis transformation with a SERK promoter fragment/luciferase gene fusion

with a CCD camera for a period of 5 times one hour (Schmidt et al. (1997) Development

124: 2049-2062). After 7 days of culture, luciferin was removed from the cultures by

extensive washing with P1-0 medium.

Wildtype WS plants were grown under standard long day conditions: 16 hours light and 8 hours dark.

The first emerging influorescense was removed in order to increase the number of influorescenses. Five days later, plants were ready for vacuum infiltration.

Agrobacterium strain C58C1 containing the transformation plasmid was grown on a LB plate with 50 mg/l kanamycin, 50 mg/l rifampicin and 25 mg/l gentamycin. A single colony was used to inoculate 500 ml of LB medium containing 50 mg/l kanamycin, 50 mg/l rifampicin and 25 mg/l gentamycin. The cultures were grown O/N at 28 degrees Celcius and the resulting log phase culture (OD600 0.8) was centrifuged to pellet the cells and resuspended in 150 ml of infiltration medium (0.5x MS medium (pH 5.7) with 5% sucrose and 10 μl/l benzylaminopurine). The inflorescenses of 6 Arabidopsis plants are submerged in the infiltration suspension while he remaining parts of the plants (which are still potted) are placed upside down on meshed wire to avoid contact with the infiltration suspension.

Vacuum is applied to the whole set-up for 10 min. at 50 kPa. Plants are directly afterwards placed under standard long day conditions. After completed seed setting the seeds were surface sterilized by a 1% sodium hypochlorite soak, then thoroughly washed with sterile water and plated onto petridishes with 0.5xMS medium and 80 mg/l kanamycin in order to select for transformed seeds. After 5 days germination under long day conditions (10.000 lux), the transformed seedlings could be identified by their green color of their cotyledons

(the untransformed seedlings turn yellow), and were further grown in soil under C1 lab conditions under long day conditions. This vacuum infiltration method resulted in approximately 0.1% transformed seeds.

Transformation of a construct containing both a gene encoding kanamycin resistance and the 2200 bp (HindIII / Dral) SERK genomic DNA fused to the firefly luciferase gene into Arabidopsis thaliana (WS) by vacuum infiltration resulted in six different kanamycin-resistent primary transformants (I, II, III, IV, V and VI). Plants IV and VI died at the seedling stage, although they were kanamycin resistant. A T2 generation could be obtained from the four plants I, II, III and V (Figure 4). Within the siliques of the T2 generation of plants no. III and V, an early inhibition in development could be observed in appoximatetely 25-50 % of the seeds. The plants I and II did not show a reduction in the number of developing seeds. (Figure 5). Similar results were observed in a T3 generation, in which again approximately 25-50% of the seeds showed an early inhibition of normal seed development.

Arabidopsis transformation with a AtSERK gene

Isolation of the AtSERK genomic and cDNA clones

Using the DcSERK cDNA sequence (seq ID no. 2) as a probe, a lambda ZipLox genomic library made form Arabidopsis Landsberg erecta total genomic DNA is screened for the presence of homologous sequences. Three different lambda clones with inserts of 14, 18 and 20 kb respectively are obtained. The 14 kb clone is digested by EcoRI and the resulting fragments subcloned into pBluescript vectors. Fragments spanning the entire coding sequence of the AtSERK gene are isolated, sequenced and compared with the Daucus homologues. The resulting sequence is shown as SEQ ID NO: 20.

Using the DcSERK cDNA sequence (SEQ ID NO: 2) as a probe, a lambda ZAPII cDNA library is screened for the presence of homologous sequences. Four lambda clones are obtained and their inserts subcloned into pBluescript vectors using the helper phage excision procedure. Fragments spanning the entire AtSERK cDNA coding sequence of the AtSERK gene are isolated, sequenced and compared with the Daucus homologues. The resulting sequence is shown as SEQ ID NO: 32.

Plasmids containing promoter sequences

Arabidopsis thaliana LTP1 promoter fragment is obtained from the binary plasmid pUH1000 (Thoma, S., Hecht, U., Kipper, A., Borella, J., De Vries, S.C., Sommerville, C. (1994) Plant Physiol. 105, 35-45) by digestion with *Bam*H1 and *HindIII* and cloning into pBluescript SK⁻ (pMT121).

- The CaMV 35S promoter enhanced by duplication of the -343 to -90 region (Kay et al., (1987) Science 236: 1299-1302) is isolated from the pMON999 vector by digestion with Hindll and Sstl and cloned into the pBluescript SK⁻ vector (pMT120).
- The promoter AtDMC1 (Klimyuk and Jones (1997) Plant Journal 11: 1-14).

 Plasmid SLJ 9691 is a construct consisting of pBluescript SK+ in which the *Arabidopsis thaliana* DMC1 genomic clone (accession number U76670) is cloned into the EcoRV site.

 SLJ 9691 carries *EcoRV* fragments of the 5' end of the AtDMC1 gene with the following modification: a BglII site instead of the second HpaI site, two ATG codons in the first exon and an Xhol site at the ATG codon of the second exon.
- The FBP7 promoter from Petunia (Angenent et al. (1995) Plant Cell 7: 1569-1582).

 The promoter of the FBP7 gene is cloned by subcloning the 0.6 kb *Hin*dIII *X*bal genomic DNA fragment of FBP7 into the *Hin*dIII *X*bal site of pBluescript KS-, resulting in the vector FBP201.

The pAtSERK binary vector constructs.

Based on the pBIN 19 vector, a binary vector pAtSERK is constructed for transformation of the *Arabidopsis thaliana* SERK cDNA under the control of different promoters.

The full length *Arabidopsis thaliana* cDNA clone of SERK (Seq ID No. NEW) is obtained from a pBluescript SK- plasmid. A Smal - KpnI 2.1 kb fragment containing the AtSERK cDNA is cloned into pBIN19 Smal - KpnI. The polyadenylation sequence from the pea rbcS::E9 gene (Millar et al., 1992), Plant Cell 4: 1075-1087) is placed downstream from the AtSERK cDNA by cloning a Klenow-filled EcoRI - HindIII E9 DNA fragment into the Klenow-filled Xmal site of the pBIN19:AtSERK vector in order to generate the binary vector pAtSERK.

Construction of plant expression vectors

The pAtSERK binary vector is used to generate the following promoter-AtSERK constructs.

- The AtLTP1 promoter is cloned in the Smal site of the pAtSERK binary vector as a Klenow-filled Kpnl-Sstl DNA fragment to give the pAtLTP1AtSERK vector.
- The CaMV 35S promoter is cloned in the Smal site of the pAtSERK binary vector as a Klenow-filled *Kpnl-Sstl* fragment to give the p35SAtSERK vector.
- The AtDMC1 promoter consisting of the Bglll Xhol 3.3kB fragment from the clone SLJ 9691 is filled in with Klenow and cloned into the Smal site of the pAtSERK binary vector to give the pAtDMC1AtSERK vector.
- A Sacl-KpnI fragment of FBP2101 is filled in with Klenow and cloned into the SmaI site of the pAtSERK binary vector to give the pFBP2101AtSERK vector.

Introduction of plant expression vectors into Arabidopsis thaliana plant cells

The above described vector constructs (pAtLTP1AtSERK, p35SAtSERK, pAtDMC1AtSERK, pFBP2101AtSERK) have been electrotransformed into *Agrobacterium tumifacienses* strain C58C1 as known in the art.

Wild type Arabidopsis thaliana WS plants are grown under standard long day conditions: 16 hours light and 8 hours dark.

The first emerging inflorescence is removed in order to increase the number of influorescences. Five days later, plants are ready for vacuum infiltration. Agrobacterium strain C58C1 containing the transformation plasmid (the pAtLTP1AtSERK vector or the p35SAtSERK or the pAtDMC1AtSERK vector or the pFBP2101AtSERK vector) is grown on a LB plate with 50 mg/l kanamycin, 50 mg/l rifampicin and 25 mg/l gentamycin. A single colony is used to inoculate 500 ml of LB medium containing 50 mg/l kanamycin, 50 mg/l rifampicin and 25 mg/l gentamycin. The cultures are grown O/N at 28 degrees Celsius and the resulting log phase culture (OD600 0.8) is centrifuged to pellet the cells and resuspended in 150 ml of infiltration medium (0.5x MS medium (pH 5.7) with 5% sucrose and 1 mg/l benzylaminopurine). The inflorescences of 6 Arabidopsis plants are submerged in the infiltration suspension while the remaining parts of the plants (which are still potted) are placed upside down on meshed wire to avoid contact with the infiltration suspension.

Vacuum is applied to the whole set-up for 10 min. at 50 kPa. Plants are directly afterwards placed under standard long day conditions. After completed seed setting the seeds are surface sterilized by a 1% sodium hypochlorite soak, then thoroughly ished with sterile water and plated onto petridishes with 0.5xMS medium and 80 mg/l kanamycin in order to select for transformed seeds. After 5 days germination under long day conditions (10.000 lux), the transformed seedlings could be identified by their green colour of their cotyledons (the untransformed seedlings turn yellow), and are further grown in soil under long day conditions. This vacuum infiltration method resulted in approximately 0.1% transformed seeds.

Expression of SERK sequences in Arabidopsis thaliana plant cells

The inflorescences from transgenic and not transgenic *Arabidopsis thaliana* plants are analysed by Whole mount *in situ* hybridisation analysis with AtSERK cDNA as probe. The inflorescences in different stages of development are fixed for 60 min. in PBS containing 70 mM EGTA, 4% paraformaldehyde, 0.25% glutaraldehyde, 0.1% Tween 20, and 10% DMSO. Samples are then washed, treated with proteinase K for 10 min, again washed and fixed a second time. Hybridisation solution consisted of PBS containing 0.1% Tween 20, 330 mM NaCl, 50 mg/ml heparin, and 50% deionized formamide. Hybridisation took place for 16 hours at 42°C using digoxigenin-labeled sense or antisense riboprobes (Boehringer Mannheim). After washing, the cells are treated with RNaseA and incubated with anti-digoxigenin-alkaline phosphatase conjugate (Boehringer Mannheim) which had been preabsorbed with a plant protein extract. Excess antibody is removed by washing followed by rinsing in staining buffer (100 mM Tris-HCl pH 9.5, 100 mM NaCl, 5 mM MgCl₂, 1 mM levamisole) and the staining reaction is performed for 16 hours in a buffer containing NBT and BCIP. Observations are performed using a Nikon Optiphot microscope equipped with Nomarski optics.

The transformed plants show ectopic expression of SERK in the vicinity of the embryo sac.

a

SPOUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT:
 - (A) NAME: NOVARTIS AG
 - (B) STREET: Schwarzwaldallee 215
 - (C) CITY: Basel
 - (E) COUNTRY: Switzerland
 - (F) POSTAL CODE (ZIP): 4058
 - (G) TELEPHONE: +41 61 69 11 11
 - (H) TELEFAX: + 41 61 696 79 76
 - (I) TELEX: 962 991
- (ii) TITLE OF INVENTION: Improvements in or relating to organic compounds
- (iii) NUMBER OF SEQUENCES: 33
- (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
- (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6695 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: unknown

- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Daucus carota
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 3696..6617
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 3731..3802
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 3851..3979
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 4124..4211
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 4284..4357
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 4430..4528
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 4642..4757

(ix) FEATURE:

(A) NAME/KEY: intron

(B) LOCATION: 4890..4967

(ix) FEATURE:

(A) NAME/KEY: intron

(B) LOCATION: 5295..5803

(ix) FEATURE:

(A) NAME/KEY: intron

(B) LOCATION: 6197..6339

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

TCTAGATGAC GAAATCGCGC TACCTTTGAT TINGAAATAC TAGGTTGTAG TATCTTG	ATT 60
AGITTITIGG ATATCITGCT GTAATTICTT TAGGAGATGC AAACGGTCTT CATTTAA	TAT 120
GAGCCCTTGT GACTTGACAA AAGTATCTAG CATGTTTGAT CACGAGGTAG CTAAAAA	GTA 180
GCGTGTTTGA TTAAGCACAT AATATTGTAT TGGGCCTATT GGCTATCAAT GAAGTTT	GAT 240
GCAAGTATAT AGCTTGTATT ATGCATGTGA TGAGGGTATA TAAAAGAAGT AAAGAAC	ATT 300
CICTOGTAGO ATTOATTITT CICTIGOCTA TAGITAACGA GTTTTGTCAC ACATGACO	FIT 360
GAAACTGGAT GTGTCTGTTC TTCCATCTAA GTTTGGATTA CCTGATAGAT GCTCAACT	PTC 420
TTCGTCAGCC TTTTCTTTCC GATTTTTCCC AAGACAAGAT TCTTTAGTTA ATAGTTAT	PTG 480
CTCTGGTGGC TTGTGTGCAT TTTAGGAATC TTACTCTGTT TTTTAATGGA GAAACGAA	NAC 540
CHACCITIT TICIGIGITC CCITITATGA TATCACCIGC TIGGAGGCGI TIAGACTI	TA 600
TCCACCTAAA CTATTCATGT TTACCAGACA AGCTATACGT TTTATCCCCC CCCCCGC	
ACCTONOGAC AAAAGAAGCG CTGATGAACT GATTTAATCC GTGTTTTATT ATATTACA	CA 720

PIGATGCITC	ATGGAGCTAA	TATCTTIGGT	TAAATTICAT	GTATATATAT	ACCCTTCCCT	780
CITICICATCG	CAGTGGCCCC	TCGITTAATT	AGCGTACTTA	ATTATCTGAT	GGATACTGTA	840
TGCTTGGCAG	ATGATGTCAT	CAGATTATAC	CAMMEMET	GCTCTACAAA	ATAAAAAAACC	900
ICTATTTATG	TICATCTITT	TGGTAACAAG	TAACTAATIG	ATGCGCTATG	TTGACAGGCG	960
ATGCATTACA	CAACITACGA	ACTAGCTTGC	AAGATCCCAA	CAATGTCCTG	CAGAGCTGGG	1020
ATCCAACCCT	TGTGAACCCT	TGCACATGGT	TICATGIGAC	ATGTAACAAT	GAAAACAGTG	1080
TTATAAGAGT	GTAGGTCACT	TCCCTTATTA	ATTTTTTAG	CAAGTTACGA	ATATTTACTC	1140
AATTGAGCAG	ATGICICITT	AAATATTTT	CTTTAATTTC	TTAGCTAAGC	GGAGCATCTA	1200
TCTTAAGTAT	CICTACTGAA	TTTAAGACAT	AATACATITT	TTAAAAAAT	CTATTAGAGT	1260
GMMMACCG	CACAGOGCAC	ATATATCTTT	TTTCTGGTAA	TICAGACAAC	CTTTCTCCCG	1320
ACGATAAAAT	' AATATAAGAT	TAACICCTIG	AACTAATTTT	TTATTTTTCT	TTTCTTTTTA	1380
TGTTCTTTGC	: AGAAAGTTTC	TTATGGICTI	TTGTGAAAAG	TACATTCTAT	GATAATTTT	1440
TGGCAACTCA	TATAAATTTA	TATATATTCC	: ATGTAGTTAT	AAGTTAAAAA	AAGCITCCTA	1500
TTAATTCCAA	GATAGAGGIT	CATTITIATA	GTTTGGGCAT	CCATGAGTTI	TTGAAAATGT	1560
CAGAAATTT	gitgagitaa	. TITTACITAC	: CAACITITAT	GGCGTCATGC	AGTGATCTTG	1620
GGAATGCAGG	ATTATCTGGT	CAAITGGITC	CICITOGCCA	GTTGAAAAAT	TTACAATACT	1680
TGTAAGACC	A TATCACTIGG	; AATGCTTTAC	TITTIATACA	, GCACAATGCT	TTCAATATCT	1740
GTTAAAAGT	G TGAAAAAGTI	GACTITCTAC	G CTTCAGCAGT	TGTTCGGATZ	ATATCTATGA	1800

AGCACTTAAA	AGGCTGGGCA	ATTTTTTTGT	TATTATTTCA	AATATTGTTA	ATTGTTACTA	1860
CTTAATATGA	TAAACTGATT	TAACTCCTCA	TGATTGGTCT	CAGTCCAATG	TGCCCTCATT	1920
AGTCACATNA	NDOFFRAAAT	GGGTTGGACA	AATATAACIT	CITITCITAA	GGTYCCAGAAA	1980
GAGCACTTAT	CAACCITGIC	TAGCGCATAA	CGTCACAGTG	GGICAGICAC	GGGCTATCCA	2040
GTTTGGGGAG	GTTTTAATGA	GCACTTATTT	ACCITGICIT	TTAAACGTCT	GAGGATGTTA	2100
TTAAAGTCTG	CATCATTCAG	ACTITAAATT	AGCACTITCA	GITGIATTAT	GAATGÓTACA	2160
TGAAAGATAC	ATATCTTAAT	GTTCCTATGC	CIGITICAAC	ATGTCTCTAA	TATTCIGITA	2220
TCTTTGTCAT	CTTAAAAATG	GCACTGATTA	AAATGTGAGA	AAGGTAGTCT	TCCAATACCA	2280
TTTCATGIAT	ACCAGAGAAT	ATCATAATTT	TTTTAAATCA	TAAGTTGGGC	CCTAGAGTTT	2340
TCTCAGTATT	GGICTATITA	TATTTTCCAC	CATTTAGAAC	TGTGTTGTCA	GATGAAAATC	2400
TTGGACTICC	ACAGAAGATC	TTATAGTAAA	AGTATICITT	AGATCTGATG	ATGAAAGTTG	2460
TCATGGTGTG	GCCTGTCCCA	GAATTTAAAT	CAATCCCATG	TCACATGTTT	GITGATCIGA	2 520
CTACTCACTG	TTAATCGAAG	AGTAACTATT	TGTGAATTAA	ATGCTTTTTT	TTTIGTTCTT	2580
CATGCTTAGC	GTTATAAAGG	TCTACGTCTG	ACTATOGITT	TTAACATGTT	ATAGETTTGT	2640
ACTGACAAGT	TTAAAGTTTC	TCTTGTTTAC	GAATTAAGAA	TATATAATAT	AAAACGCTTT	2700
AACTITICICT	GTGGAAGGTG	TICTTACCIT	TTTATATATA	TATATAGATA	CTCAGACTCT	2760
GCTGGCAATT	ATATCTTACG	AACTTACGAG	TATACAGAAC	TIGTATATTA	GGTTCAGATG	2820
AGTGGCTGTA	GTAGAACACC	TTAAGCAAGA	ACTTAATCAT	GAGGITTCAA	CCTTTTAACT	2880
TICITTITAG	ATTTTTTCAA	GITTATGGAA	AATTGTACCT	CATGATCGTG	GPTTCTTTCC	2940

ATAAACTTTC	CATATAAGTC	CGTTTCTTGA	CGTTTTCATG	TAAGCTGTTG	ACGAGTGATT	3000
ATTAGCGGIT	CTTTCAATAA	TCATAATGIG	TCTCACTTIG	ATGAGGCCTG	TACTTATTAT	3060
TGCACCTTGC	ACTTAACCTT	GATCCTCATG	TCATCTTGAT	TGICATAGIC	TACTAACCGA	3120
GTTGAACATG	GTTTATCATG	TCTTTTGAGG	TAACAATGTA	GCTTTCACCT	CIGICCITGA	3180
TATAGGITTA	AGGCTTGCAC	CTCCCACTAG	cerricerie	TTTTATTCAC	AGTTCACACA	3240
CCTACTAGCA	CIGITCACCI	CTAGICTTT	GTCCGCAAAT	agtaagaagt	TICITICGCA	3300
TAATAGTGGA	TGATCATTTA	AGAAATAGTG	AATCAAATTA	TOGIGITATT	GIGITIGIAC	3360
TTTGGAATTA	AATGAGTTGC	TGAACATTGT	TGCTGTTTAT	CGTTGTCAAG	GCTTTGCCAA	3420
GGAAGGCGAT	TAGTAAGAGT	GGGCATCCAA	GCGCCTTTAT	CTTGAAGGGG	CGGGCGGCAC	3480
GIIGIGGATT	CIGGGIGICI	ATTAGAGGAC	ATTATCTATA	TATACTGATT	ATTTATTAGA	3540
ATATAAATCA	ACTACTATAT	TTTTCTTTGT	AATGITTATA	TAGAAATCCC	ACTCGTAAAC	3600
TTGACAAATA	CCATTGAAAT	ATTIGAACCI	AATTAATTAG	TAGIGICAGG	TITAAATICA	3660
AACTCATTTA	ATTTTACTTT	ААААААТААТ	TCTATATGAA	TCGTAACAGT	ATAAATATAT	3720
TAAATTACAT	GTATGTGTGC	CTATATATAG	CIGAATGICI	AATAGACTCC	AAGACGGCTG	3780
CICITACIGO	CTAGGCGTCC	AGGCAGTTCA	CTGATGCTTA	CCTTGACAAA	TATGGGGTTC	3840
GTATGACATT	GPTGGGGATC	CCTATCACTG	GATTCCIGIT	TTGCTGACCC	TCTGTTCAAT	3900
TGATITICAT	TGATGTAGTA	TTACTAGTIT	TATAAATATT	CTTTATTGCA	ATAATITAAC	3960
TGGAGTTTAA	CAATGACAGG	GAGCTTTACA	GCAATAACAT	AAGTGGACCA	ATTCCTAGTG	4020

ATCTTGGGAA TCTGACAAAT TTGGTGAGCT TGGACCTATA CATGAATAGC TTCTCTGGAC	4080
CTATACCGGA CACATTAGGA AAGCTTACAA GGCTAAGATT CITGTATGAC TACAAATCTT	4140
CACTAGITIT TAACTTAATG CAATTTGATT ATCCTTTCAA GTGATTGATT ATATCACAAA	4200
TTACIGGATA GGCGTCTCAA CAACAACTGC CTCTCTGGTC CAATTCCAAT GTCACTGACT	4260
AATATTACAA CICITCAAGT CCIGTAAGTA TICCGACCIT TCCAGATAGT TITGITGITG	4320
TGGATGITTC AATTITAATA CTAAATATGT TCATCAGGGA TITATCAAAC AATCGGCTAT	4380
CAGGACCAGT ACCGGATAAT GGCTCATTTT CTTTGTTTAC ACCTATCAGG TTTAATGCTA	4440
GTAATATCTT TAATATTATG GTTCTTACTT CTACTGCGAA AGCTATGATA ATATTTTTTT	4500
TCTCCTTCAT ATATTATCAC TTTCGCAGTT TTGGCAATAA TTTGAATTTA TGTGGACCTG	4560
TAACTGGGAG GCCCTGCCCT GGATCTCCCC CATTTTCTCC ACCACCTCCG TTCATCCCAC	4620
CATCAACAGT ACAGCCTCCA GGTGATTTAG TTTTTATATT AATTCCCGTA ATTAATTTTA	4680
TGACTGTAAA AATTGGTGTT AATTTCACCA GTTGCGAATA AAGTATTTTC CTTCTTTCTC	4740
TICITATTAT TATGAAGGAC AAAATGGTCC CACTGGAGCT ATTGCTGGGG GAGTAGCTGC	4800
TOGTGCTGCT TTACTGTTTG CTGCACCTGC AATGGCATTT GCATGGTGGC GGAGAAGAAA	4860
ACCGCGAGAA CATTICITIG ATGIGCCAGG TTAGICCTGT AAATAGATAT CTATTGAAGC	4920
GCTTACTGTC TGTGGACTTT GTTTTCACTG TCATTAGTTA ACTTCAGCTG AAGAGGACCC	4980
AGAAGTGCAC CTTGGTCAAC TGAAGAGGTT TTCTCTGCGA GAATTGCAAG TCGCAACGGA	5040
TACTITITAGT ACCATCCITG GAAGAGGIGG ATTIGGTAAG GIGIATAAGG GACGCCITGC	5100
TGATGGCTCA CTTGTAGCAG TTAAAAGGCT TAAAGAAGAA CGAACACCAG GTGGCGAGCT	5160

PCT/EP97/02443

GCAGITICAA ACAGAAGIGG AAAIGATTAG CATGGCIGIG CATCGAAAIC TICIGCGICT 5220 ACGIGGITTC TGCATGACAC CTACCGAGCG GCTTCTTGTA TATCCATACA TGGCTAATGG 5280 AAGTGTTGCG TCATGTTTAA GAGGTATCTC AGTTACAATT ACCATAACTT GCCAGAAGTT TGITTGATTA AAAATGAAAT ATAACTCCCT ACACTATGIT AAGGIGITAT AATTTCIGAG 5400 CAGATCTTAT TTCCCATTGC AAGATACCAG TTATTATTGT TTTTTCTGTA ATTGATACCG 5460 GITATATTIC TITCIIGTAT TIGGITATAT GCAAGGATTI CGAGICTAAT AAGITATCAA 5520 ACTGGATGCT ATGTTTATTC TGCAATTGAA TTCTTGCTTC ATGTGCCAAA ATATATATGA 5580 TTCAACTTGG AATCATCTTA TAATATACTG TGTAAAGTCA GCTGTTGACT TTCATCATTA 5640 ATTAGTOTTC ATAAATCAGA ATCTGCCTAG TGAGCTTTAC CGACATACTC TAAACCTTTC 5700 TTATGGCCCT GTATATAATC GTCCCACTTA CTTTATTCAG TITGTCTGCT CTCTGAATTT 5760 TTGATCTGTA CATTGTGATG TCTTGTTTTC ATCAAATGTA GAGCGTCAGC CATCAGAACC 5820 TCCCCCTGAT TGGCCAACTA GGGAGAGGAT TGCACTAGGA TCTTCTAGGG GCCTATCTAA 5880 ATTGCATGAC CATTGTGATC CCAAGATTAT CCATCGCGAT GTAAAAGCTG CAAATATATT 5940 6000 ATTGGACGAA GAATTTGAGG CTGTTGTAGG TGATTTTGGG TTAGCTAGGC TCATGGATTA CAAGGATACC CATGITACGA CTGCTGTAAG GGGTACCATT GGGCACATAG CTCCCGAGTA 6060 CCTCTCGACT GGAAAGTCAT CAGAGAAGAC CGATGTCTTT GGTTATGGGA TAATGCTCCT 6120 AGAGCTCATT ACTGGACAGA GGGCTTTTGA TCTTGCTCGC CTTGCGAACG ATGATGATGT 6180 6240 TATGTTGTTG GATTGGGTAT GTGTCCCGGG TGTTCCTTTG GTTAATTATT TCACATATTA

GIGCITACTA	CTTIGTTGTG	GCCCTTTGTT	TPTATTICCT	GCCIGTATTT	CATTCTTAGT	6300
CATGITIATGO	ATATIGACCT	GCTTTGCAAT	GTCTTTTAGG	TTAAAAGCCT	TTTGAAAGAG	6360
AAAAAGTTGG	AGATGCTGGT	CGATCCTGAC	CTGCAGAACA	ATTACATTGA	CACAGAAGTT	6420
GAGCAGCTTA	TTCAAGTAGC	ATTACTCTGT	ACCCAGGGTT	CGCCAATGGA	GCGGCCTAAG	6480
ATGTCAGAGG	TAGTCCGAAT	GCTTGAAGGT	GATGGCCTTG	CAGAAAAGTG	GGACGAGTGG	6540
CAAAAAGTTG	AAGTCATCCA	TCAAGACGTA	GAATTAGCTC	CACATOGAAC	TICIGAATGG	6600
ATCCTAGACT	CGACAGATAA	CTTGCATGCT	TTTGAATTAT	CTGGTCCAAG	ATAAACAGCA	6660
TATAAAATGT	AATGAAATTA .	ATATTTTTTA	TCCTT			6695

(2) INFORMATION FOR SEQ ID NO: 2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1815 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEINESS: single
 - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: CDNA
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Daucus carota
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 94..1752

(xi)	SECUENCE	DESCRIPTION:	SEO	ID	NO:	2:	
------	----------	--------------	-----	----	-----	----	--

GACAAATACC ATTGAAATAT TIGAACCTAA TTAATTAGTA GIGTCAGGTT TAAATTCAAA	. 60
CTCATTTAAT TTTACTTTAA AAAATAATTC TAT ATG AAT CGT AAC AGT ATA AAT Met Asn Arg Asn Ser Ile Asn 1 5	114
ATA TTA AAT TAC ATG CAG TTC ACT GAT GCT TAC CTT GAC AAA TAT GGG Ile Leu Asn Tyr Met Gln Phe Thr Asp Ala Tyr Leu Asp Lys Tyr Gly 10 15 20	162
GTT CTT ATG ACA TTG GAG CTT TAC AGC AAT AAC ATA AGT GGA CCA ATT Val Leu Met Thr Leu Glu Leu Tyr Ser Asn Asn Ile Ser Gly Pro Ile 25 30 35	210
CCT AGT GAT CTT GGG AAT CTG ACA AAT TTG GTG AGC TTG GAC CTA TAC Pro Ser Asp Leu Gly Asn Leu Thr Asn Leu Val Ser Leu Asp Leu Tyr 40 45 50 55	258
ATG AAT AGC TTC TCT GGA CCT ATA CCG GAC ACA TTA GGA AAG CTT ACA Met Asn Ser Phe Ser Gly Pro Ile Pro Asp Thr Leu Gly Lys Leu Thr 60 65 70	306
AGG CTA AGA TTC TTG CGT CTC AAC AAC AAC AGC CTC TCT GGT CCA ATT Arg Leu Arg Phe Leu Arg Leu Asn Asn Asn Ser Leu Ser Gly Pro Ile 75 80 85	354
CCA ATG TCA CTG ACT AAT ATT ACA ACT CTT CAA GTC CTG GAT TTA TCA Pro Met Ser Leu Thr Asn Ile Thr Thr Leu Gln Val Leu Asp Leu Ser 90 95 100	402
AAC AAT CGG CTA TCA GGA CCA GTA CCG GAT AAT GGC TCA TIT TCT TTG Asn Asn Arg Leu Ser Gly Pro Val Pro Asp Asn Gly Ser Phe Ser Leu 105 110 115	450
TIT ACA CCT ATC AGT TIT GCC AAT AAT TIG AAT TIA TGI GGA CCC GTA	498

WO 97/43427 PCT/EP97/02443

		r P	ro	Ile	e Se	r Ph	e Al	a As	n As	n Le	u Ası	n Le	u Cyr	s Gl	y Pr	o Va	1
120)					12	5				130	0				13	5
ACT	GG	G A	GG	α	TG	: cc	rec	A TC	r cc	c ec	A TT	rio	s ccz	A CC	A CC	T CO	G 546
											o Phe						
					140					145	_				15		_
											A GGA						
Phe	Ile	≥ P1	0			Th	· Val	Glr	Pro	Pro	Gly	Gl	ı Asr	Gly	Pro	o Thi	•
				155					160)				165	5		
GGA	G(~	ר ב	T)	ىتت	ccc												
											GCT						
2		17			GIY	GLY	Val	. A18		GIY	Ala	Ala			Phe	≥ Ala	l
			_					1/3	ı				180				
GCA	ccı	. GC	Ά	ATG	GCA	TTT	GCA	TGG	TGG	CCC	AGA	AGA	444	CCC:	. (722	ממם י	600
											Arg						
	185						190		_	_	_	195					•
											CCA						
	Phe	Ph	e .	Asp	Val	Pro	Ala	Glu	Glu	Asp	Pro	Glu	Val	His	Leu	Gly	
200						205					210					215	
CNA	~~~																
CAA																	786
Gln	Deu	Ly	5 &			ser	Leu	Arg	Glu		Gln	Val	Ala	Thr		Thr	
					220					225					230		
TTT	AGT	AC	C #	ATA	CIT	GGA	AGA	GGT	GGA	بلمليد	GGT	AAC	CALC.	ጥአጠ	220	CCA	07.4
Phe	Ser	Thi	- 1														834
				:35		_		-	240		1	-,-	· · ·	245	Lys	GIY	
ccc	CIT	GC	י פ	AT (œc	TCA	CTT	GTA	GCA	GTT	AAA	AGG	CTT	AAA	GAA	GAA	882
Arg :	Leu	Ala	ı A	sp (Gly	Ser	Leu	Val	Ala	Val	Lys	Arg	Leu	Lys	Glu	Glu	-
		250						255					260				
CGA 2																	930
Arg !	lhr	Pro	G	ly (Sly	Glu	Leu	Gln	Phe	Gln	Thr	Glu	Val	Glu	Met	Ile	

	265					270					275					
»CC	באדע	للمك	GTG	САТ	CGA	TAA	CIT	crg	CGT	CTA	CGT	GGT	TTC	TGC	ATG	978
			Val													
280	Mec	,	·		285	-				290					295	
200																
ACA	CCA	ACA	GAG	CGG	CTT	CTT	GTA	TAT	CCA	TAC	ATG	GCT	TAA	GGA	AGT	1026
			Glu													
				300					305					310		
GTT	GCG	TCG	TGT	TTA	AGA	GAG	CCI	CAG	CCA	TCA	GAA	CCT	ccc	CTT	GAT	1074
Val	Ala	Ser	Cys	Leu	Arg	Glu	Arg	Gln	Pro	Ser	Glu	Pro	Pro	Leu	Asp	
			315					320					325			
																1122
			AGG													1122
Trp	Pro	Thr	Arg	Lys	Arg	Ile	Ala	Leu	Gly	Ser	Ala		Gly	Leu	Ser	
		330					335					340				
										.	C. M.	~~	CNT	עדיא	מממ	1170
			GAC													1170
Tyr			Asp	His	Cys			гуs	TIE	116	355		MSP	Val	Lys	
	345					350					درد					
COM	003	አንጥ	' ATA	עיודי	ant.	GAC	CAA	CAA	بلملمان	GAG	GCT	r GTT	GTA	GGT	GAT	1218
			Ile													
360		ASI	11-	Deu	365		0_0			370				-	375	
300					505											
TTT	GGG	TTA	GCI	AGG	CTC	ATG	GAT	TAC	AAG	GAT	, VCC	CAT	GII	ACA	ACT	1266
			ı Ala													
	_			380					385					390		
GCT	CIP	AGC	GGI	ACC	TTG	GGC	TAC	ATA	CCI	. 000	GAC	TAC	CTC	TCG	ACT	1314
Ala	. Val	. Arg	g Gly	Thr	Leu	Gly	Tyr	: Ile	Ala	Pro	Gli	ı Tyr	Leu	Ser	Thr	
			395	5	٠			400)				405	i		
															CIC	1362
Gly	Lys	s Sec	c Sez	Glu	ı Lys	Thu	Asp	val	l Phe	e Gly	Ty			Met	Leu	
		41	0				415	5				420)			

TTA	GAG	CIC	ATT	ACT	GGA	CAG	AGA	GCT	TTT	GAT	CTT	GCT	CCC	CTT	GCG	1	410
Leu	Glu	Leu	Ile	Thr	Gly	Gln	Arg	Ala	Phe	Asp	Leu	Ala	Arg	Leu	Ala		
	425					430					435						
AAC	GAT	GAT	GAT	GTT	ATG	TIG	TIG	GAT	TCG	GIT	AAA	AGC	CIT	TIG	AAA	1	458
Asn	Asp	Asp	Asp	Val	Met	Leu	Leu	Asp	Trp	Val	Lys	Ser	Leu	Leu	Lys		
440					445					4 50					455		
GAG	AAA	AAG	TTG	GAG	ATG	CIG	GIC	GAT	CCI	GAC	CIG	GAG	AAC	TAA	TAC	1	.506
Glu	Lys	Lys	Leu	Glu	Met	Leu	Val	Asp	Pro	Asp	Leu	Glu	Asn	Asn	Tyr		
				460					465					470			
ATT	GAC	ACA	GAA	GIT	GAG	CAG	CIT	ATT	CAA	GTA	GCA	TTA	crc	TGT	ACC	1	.554
Ile	Asp	Thr	Glu	Val	Glu	Gln	Leu	Ile	Gln	Val	Ala	Leu	Leu	Cys	Thr		
			475					480					485				
CAG	GGT	TCG	CCA	ATG	GAG	ccc	CCI	AAG	ATG	TCA	GAG	GTA	GIC	CGA	ATG	1	.602
Gln	Gly		Pro	Met	Glu	Arg	Pro	Lys	Met	Ser	Glu	Val	Val	Arg	Met		
		490			•	٠.	495					500					
	GAA															1	.650
Leu	Glu	GIY	Asp	Gly	Leu		Glu	Lys	Trp	Asp		Trp	Gln	Lys	Val		
	505					510					515						
<i>~</i>	~~~		~~ m		~~~	~~~	~~~			201	~~ m				~		600
	GIC															1	.698
	Val	TIE	HIS	GIN		Val	GIU	Leu	Ala		HIS	Arg	Inr	ser			
520					525					530					535		
TYCE:	ATC	עידי	CNC	m~	מרים	CMT	አአሮ	mm-	Chm	درس	بلغلفان	CAR	OTT DA	سمي	~~	1	746
	Ile															1	. / 140
			-wp	540	***	nap	WO(1	Deu	545	AL a	21165	014	Deu	550	GIJ		
				J-#U					J#J					J.0			
CCA	AGA	TAA	ACAC	CAT :	ΑΤΑΑ	AATC	IG A	ATGA	AATT	A AT	ململمك	מייוין	MX:	TTAA:	AAA	1	802
_	Arg				m M									: = 1		•	
	9																

AAAAAAAAA AAA

- (2) INFORMATION FOR SEQ ID NO: 3:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 553 amino acids

(B) TYPE: amino acid(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

Met Asn Arg Asn Ser Ile Asn Ile Leu Asn Tyr Met Gln Phe Thr Asp 1 5 10 15

Ala Tyr Leu Asp Lys Tyr Gly Val Leu Met Thr Leu Glu Leu Tyr Ser 20 25 30

Asn Asn Ile Ser Gly Pro Ile Pro Ser Asp Leu Gly Asn Leu Thr Asn 35 40 45

Leu Val Ser Leu Asp Leu Tyr Met Asn Ser Phe Ser Gly Pro Ile Pro 50 55 60

Asp Thr Leu Gly Lys Leu Thr Arg Leu Arg Phe Leu Arg Leu Asn Asn 65 70 75 80

Asn Ser Leu Ser Gly Pro Ile Pro Met Ser Leu Thr Asn Ile Thr Thr 85 90 95

Leu Gln Val Leu Asp Leu Ser Asn Asn Arg Leu Ser Gly Pro Val Pro
100 105 110

Asp Asn Gly Ser Phe Ser Leu Phe Thr Pro Ile Ser Phe Ala Asn Asn 115 120 125 Leu Asn Leu Cys Gly Pro Val Thr Gly Arg Pro Cys Pro Gly Ser Pro

Pro Phe Ser Pro Pro Pro Pro Phe Ile Pro Pro Ser Thr Val Gln Pro

Pro Gly Gln Asn Gly Pro Thr Gly Ala Ile Ala Gly Gly Val Ala Ala

Gly Ala Ala Leu Leu Phe Ala Ala Pro Ala Met Ala Phe Ala Trp Trp

Arg Arg Arg Lys Pro Arg Glu His Phe Phe Asp Val Pro Ala Glu Glu

Asp Pro Glu Val His Leu Gly Gln Leu Lys Arg Phe Ser Leu Arg Glu

Leu Gln Val Ala Thr Asp Thr Phe Ser Thr Ile Leu Gly Arg Gly Gly

Phe Gly Lys Val Tyr Lys Gly Arg Leu Ala Asp Gly Ser Leu Val Ala

Val Lys Arg Leu Lys Glu Glu Arg Thr Pro Gly Gly Glu Leu Gln Phe

Gln Thr Glu Val Glu Met Ile Ser Met Ala Val His Arg Asn Leu Leu

Arg Leu Arg Gly Phe Cys Met Thr Pro Thr Glu Arg Leu Leu Val Tyr

Pro Tyr Met Ala Asn Gly Ser Val Ala Ser Cys Leu Arg Glu Arg Gln

Pro Ser Glu Pro Pro Leu Asp Trp Pro Thr Arg Lys Arg Ile Ala Leu

				325					330					335	
Gly	Ser	Ala	Arg 340	Gly	Leu	Ser	Tyr	Leu 345	His	Asp	His	Cys	As p 350	Pro	Lys
Ile	Ile	His 355	Arg	Asp	Val	Lys	Ala 360	Ala	Asn	Ile	Leu	Leu 365	Asp	Glu	Glu
Phe	Glu 370	Ala	Val	Val	Gly	Asp 375	Phe	Gly	Leu	Ala	A rg 380	Leu	Met	Asp	Tyr
Lys 385	Asp	Thr	His	Val	Thr 390	Thr	Ala	Val	Arg	Gly 395	Thr	Leu	Gly	Tyr	Ile 400
Ala	Pro	Glu	Tyr	Leu 405	Ser	Thr	Gly	Lys	Ser 410	Ser	Glu	Lys	Thr	Asp 415	Val
Phe	Gly	Tyr	Gly 420	Ile	Met	Leu	Leu	Glu 42 5	Leu	Ile	Thr	Gly	Gln 4 30	Arg	Ala
Phe	Asp	Leu 435	Ala	Arg	Leu	Ala	Asn 440	Asp	Asp	Asp	Val	Met 445	Leu	Leu	Asp
Trp	Val 450	Lys	Ser	Leu	Leu	Lys 4 55	Glu	Lys	Lys	Leu	Glu 460	Met	Leu	Val	Asp
Pro 465	Asp	Leu	Glu	Asn	Asn 470	Tyr	Ile	Asp	Thr	Glu 475	Val	Glu	Gln	Leu	Ile 480
Gln	Val	Ala	Leu	Leu 485	Cys	Thr	Gln	Gly	Ser 490		Met	Glu	Arg	Pro 495	Lys
Met	Ser	Glu	Val 500		Arg	Met	Leu	Glu 505		Asp	Gly	Leu	Ala 510	Glu	Lys
Trp	Asp	Glu	Trp	Gln	Lys	Val	Glu	Val	Ile	His	Gln	Asp	Val	Glu	Leu

520

. 525

515

WO 97/43427 PCT/EP97/02443 - 54 -

Ala Pro His Arg Thr Ser Glu Trp Ile Leu Asp Ser Thr Asp Asn Leu 530 535 540

His Ala Phe Glu Leu Ser Gly Pro Arg 545 550

- (2) INFORMATION FOR SEQ ID NO: 4:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 13 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: primer
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

TETTETET TGC 13

- (2) INFORMATION FOR SEQ ID NO: 5:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 10 base pairs

(B) TYPE: nucleic acid

(C) STRANDEINESS: single

(D) TOPOLOGY: unknown

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: primer
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

GGGATCTAAG 10

- (2) INFORMATION FOR SEQ ID NO: 6:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 10 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: primer
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

ACACGIGGIC 10

- (2) INFORMATION FOR SEQ ID NO: 7:
 - (i) SECUENCE CHARACTERISTICS:

(A) LENGIH: 10 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

(iii) HYPOTHETICAL: NO

- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:

(A) ORGANISM: primer

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

TCAGCACAGG

10

- (2) INFORMATION FOR SEQ ID NO: 8:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 14 base pairs

(B) TYPE: nucleic acid

- (C) STRANDEDNESS: single
- (D) TOPOLOGY: unknown
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: primer
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

TITTITITIT TCIG

14

- (2) INFORMATION FOR SEQ ID NO: 9:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 13 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

(iii) HYPOTHETICAL: NO

- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: primer
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

TTTTTTTTT TCA

- (2) INFORMATION FOR SEQ ID NO: 10:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDELNESS: single
 - (D) TOPOLOGY: unknown
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: primer
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

GACATCGTCC 10

- (2) INFORMATION FOR SEQ ID NO: 11:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: unknown

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: primer
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

CCCTACTGGT 10

- (2) INFORMATION FOR SEQ ID NO: 12:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDETNESS: single
 - (D) TOPOLOGY: unknown
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: primer
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

ACACGTGGTC 10

- (2) INFORMATION FOR SEQ ID NO: 13:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: unknown

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE:

(A) ORGANISM: primer

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

GGTGACTGTC 10

- (2) INFORMATION FOR SEQ ID NO: 14:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 18 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:

(A) ORGANISM: primer

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

TCTTGGACCA GATAATTC

- (2) INFORMATION FOR SEQ ID NO: 15:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 19 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE:

(A) ORGANISM: primer

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

CICIGATGAC TITICCAGIC

19

- (2) INFORMATION FOR SEQ ID NO: 16:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 16 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE:

(A) ORGANISM: primer

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

AATGGCATTT GCATGG

16

- (2) INFORMATION FOR SEQ ID NO: 17:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 5 amino acids

(B) TYPE: amino acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

(ii) MOLECULE TYPE: peptide

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Daucus carota

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

Ser Pro Pro Pro Pro

- (2) INFORMATION FOR SEQ ID NO: 18:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 8 amino acids

(B) TYPE: amino acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: unknown

- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:

(A) ORGANISM: Daucus carota

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:

His Arg Asp Val Lys Ala Ala Asn

1 5

- (2) INFORMATION FOR SEQ ID NO: 19:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 9 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: unknown
 - (ii) MOLECULE TYPE: peptide
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Daucus carota
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:
 - Gly Thr Leu Gly Tyr Ile Ala Pro Glu 1 5
- (2) INFORMATION FOR SEQ ID NO: 20:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 4081 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (iii) HYPOTHETICAL: NO
 - (vi) ORIGINAL SOURCE:

(A) ORGANISM: Arabidopsis thaliana

(vii) IMMEDIATE SOURCE:

(B) CLONE: Arabidopsis SERK gene

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 1280..1367

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 1796..1928

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 2014..2085

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 2203..2346

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 2450..2521

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 2617..2688

(ix) FEATURE:

(A) NAME/KEY: excon

(B) LOCATION: 2772..2884

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 3015..3146

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 3305..3646

(ix) FEATURE:

(A) NAME/KEY: excon

(B) LOCATION: 3760..4081

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:

TCTAGAAACC	TTTTGATCAT	AATGAAAATA	AAGAGTCCAT	CCACCACATG	GGGTAAGCAT	60
AATGIGIGAI	ATTTAAAGGG	TAACAAATGT	AATCIGCITT	TTATTTTACT	TTTTACCTCT	120
ACTCAAATTG	TATGGGCAGT	TTTTTTTTT	TTTTAAATGA	TAAGACAAGT	ATCTGTTTAA	180
TGGTATTGTG	ATGAAACAGT	AGIAAAGICA	TATCGGGCAC	GCCATACTAC	TTCCACAGTG	240
GAACTTGGCC	AAATTTTGTC	TTTGCCGTCT	CTACAGTTTC	TTCCACCAAA	TTTTTTGTTG	300
ACAAAACTCA	AATCTTTCAA	TCTCATCTCT	GCCAAAGTTG	GGTTTAGAAA	GAATATCAGC	360
AAACACTAAT	ATCTTTATTG	TIGCATGGIT	TATCAATCAC	AAAATTCACA	ACCATTGTAA	420
AAAAAATTC	ACATTTTTGG	TATGAGATTG	CTCACATGAT	AGTGAACCTC	TTTAACATTT	480
TAACTTTACT	TICATAAATA	CGGGATTACG	AATCITACIT	GCATTAAAAA	TTTAGAAAAG	540
GITTITCTAC	TTAAAGAAAA	AAGGGACCCA	ACAGAGAGAG	GTTTGACCAG	GAGAAACGGG	600
TGCATAGCCT	TAAGAGCTTT	CAACTACTTT	ACCCCAAACC	CAAAGCGATG	TCACTITCAA	660
CCATCTCTTC	TCTCCCCCCA	ACCCGTTTTT	TTGACCGGTC	AGTTCGGGCA	GCAGCACCGT	720
TACGGGCAGC	TTATATTCCT	carerrocre	CTCTACACCA	CTGCATGCCC	ATAAATAAAG	780

WO 97/43427 PCT/EP97/02443

CCCGITGAGA	TCTTTAAAAA	TATTAAATAA	TATATCAACG	AAAAAGCTAT	THATICATA	840
AGAAGAAAAA	GAGAGGAACA	ACAACAACAC	ACTAATCATA	GILICICICE	CAGGCTTGTT	900
GITGCGGCTT	AATAAAAAGC	TCTTTTGTTA	TTATTACTTC	ACGIAGATIT	TCCCCAAAAA	960
GCTCTTATTT	TTTTGTTTAA	AAAAAAAGT	TTCATCTTTA	TICAACTITT	GITITACAGT	1020
GIGIGIGIGA	GAGAGAGAGT	GIGGITIGAT	TGAGGAAAGA	CGACGACGAG	AACGCCGGAG	1080
AATTAGGATT	TTTATTTTAT	TTTTTACTCT	TIGITIGITT	TAATGCTAAT	GGGITTITAA	1140
AAGGGTTATC	GAAAAAATGA	GIGAGITIGI	GTTGAGGTTG	TCTCTGTAAA	GIGITAATGG	1200
TGGTGATTTT	CGGAAGITAG	GETTTTCTCG	GATCTGAAGA	GATCAAATCA	AGATTCGAAA	1260
TTTAGCATTG	TTGTTTGAAA	TGGAGTCGAG	TTATGIGGIG	TTTATCTTAC	TITCACTGAT	1320
CITACITCCG	AATCATTCAC	TGTGGCTTGC	TICIGCTAAT	TIGGAAGGIT	CGTGGTTACT	1380
CAATTACTCA	GCTTTACTCG	TTTCTCAATT	ACTITICICGA	TICTTTTTA	TTTGGAGGTG	1440
AATCGCTATC	TTIAGIGICI	GCATTTTGAT	TTATGAAAAT	TGTTGTTGTT	CTTTGTATTT	1500
GTAAGATTTA	GTGGCTAGTA	CTTTGAATAC	ACIGITITICO	TTTTCTTGTT	CAGATCAACT	1560
TIGIATATIG	TAAAGGCATG	THETHIGGE	TGAAAAGCTG	GGTTATTTGA	TATCTTAAGA	1620
TIGATGITGT	TGATCCAAAC	ATTCTCTGAA	AGACTTCATT	TGTTTTTGGT	TTTGTAAAGA	1680
ATTIGITTAA	TRATTAGCCT	CTAATCTCAG	ACACCCTCT	TIGAATAGIT	CTCTCTTGAA	1740
ATTAGACTTT	TCACCAATTG	ATGCTAATTG	TGTAGATTTG	TIGITCITGI	TATAGGTGAT	1800
GCTTTGCATA	CTTTGAGGGT	TACTCTAGIT	GATOCAAACA	ATGTCTTGCA	GAGCTGGGAT	1860
CCTACGCTAG	TGAATCCTTG	CACATGGTTC	CATGTCACTT	GCAACAACGA	GAACAGTGTC	1920

ATAAGAGTGT	AAAGCTTTCT	TCTACTAATC	CCACTTTTTA	AACTTTGACC	TCAGCGTGGT	1980
TACCGACATT	TEIGEFFICIT	TTGTCAAATA	CACTGATTTG	GGGAATGCAG	AGITATCTOG	2040
CCATTIAGTT	CCAGAGCTTG	GTGTGCTCAA	GAATTTGCAG	TATTTGTAAG	TTCCACTTAT	2100
GCATCATGCT	TTAACAAAAC	AAATCCAAGA	TTTGACAGAA	GAAGCACTGG	AGTTACCTTT	2160
TGTAATTGAA	ATCTTTTAA	CAAGTTTCTT	ATTTTCTTAC	AGGGAGCTTT	ACAGTAACAA	2220
CATAACTGGC	CCGATICCIA	GTAATCTTGG	AAATCTGACA	AACTTAGTGA	GTTTGGATCT	2280
TTACTTAAAC	AGCTTCTCCG	GICCTATICC	GGAATCATTG	GGAAAGCTTT	CAAAGCTGAG	2340
ATTICICIGA	GTATACATAT	GCTTTACCGG	CTCAGTTACA	GICTTIGTT	AATCTTAGGT	2400
TTTGTTCCAA	TTTTTGACTC	TTTGCTGAAA	ATTITACATG	CAAGAATAGC	CGGCTTAACA	2460
ACAACAGTCT	CACTGGGTCA	ATTCCTATGT	CACTGACCAA	TATTACTACC	CTTCAAGIGI	2 52 0
TGTGAGTCCT	CTCATTAACT	TTCATTTATG	TCTACTTCAT	TCTCCCTCAG	TTGATTTGTT	2580
GAGITAATGC	ACTTAACCTT	GATGGATGCA	ACACAGAGAT	CTATCAAATA	ACAGACTOTO	2640
TOGITCAGIT	CCTGACAATG	GCICCITCIC	ACTOTICACA	CCCATCAGGT	TCTATGATTT	2700
ATCCTCTTCA	GTTATTTCAG	TIGHTGIGIC	AGIGICIGAA	CITATICIGA	AACTTTCATT	2760
TCCTTGTGCA	GITTIGCTAA	TAACTTAGAC	CTATGTGGAC	CIGITACAAG	TCACCCATGT	2820
CCTGGATCTC	CCCCGTTTTC	TCCTCCACCA	CCTTTTATTC	AACCTCCCCC	AGTITICCACC	2880
CCGAGTAAGC	CTCCTCTTT	TAGITTACAT	TATAGGAAAC	AGAAGATGAA	ATCTTTGCTT	2940
CICIGICAAT	cerrrrere	ATATAACTCA	TCTTGCCAAT	AAGGCAATAA	CCAAATGATC	3000

TAATTIGATT TCAGGIGGGT ATGGIATAAC	TGGAGCAATA	GCTGGTGGAG	TIGCIGCAGG	3060
TECTECTTIC CTCTTTCCTC CTCCTCCAAT	AGCCTTTGCT	TGGTGGCGAC	GAAGAAAGCC	3120
ACTAGATATT TTCTTCGATG TGCCTGGTGA	GITTATTATT	CGCATTAGIT	TCIGITCTIA	3180
GCCAGCAATT TTGTTTTGCA GAAAAGTATT	GGAACAACTG	TTAATGAAAA	TCAATACATA	3240
AGTCATTGTT TTTTAAGTTA CAAACTCTTT	'TGAGTAAAAT	CTCGATTGCA	AAATCTCTAT	3300
GCAGCCGAAG AAGATCCAGA AGTTCATCTG	GGACAGCTCA	AGAGGTTTTC	TTTGCGGGAG	3360
CTACAAGIGG CGAGIGAIGG GITTAGIAAC	AAGAACATTT	TGGGCAGAGG	TEEGITTEGG	3420
AAAGTCTACA AGGGACGCTT GGCAGACGGA	. ACTOTTGTTG	CTGTCAAGAG	ACTGAAGGAA	3480
GAGCGAACTC CAGGTGGAGA GCTCCAGTTT	CAAACAGAAG	TAGAGATGAT	AAGTATGGCA	3540
GTTCATCGAA ACCTGTTGAG ATTACGAGGT	TICIGIATGA	CACCGACCGA	GAGATIGCTT	3600
GIGIATOCIT ACATGGOCAA TOGAAGIGII	GCINCAIGIC	TCAGAGGTAA	AAACTAAACA	3660
ATTAAACATC TIGIGCICIC TCICAATTAC	TTTGACGTGA	AGIGITITT	CATGITITCC	3720
TITATGGGTT CATAATTGTT GGTTACACTA	. ATGACACAGA	GAGGCCACCG	TCACAACCTC	3 78 0
CGCTTGATTG GCCAACGCGG AAGAGAATCG	CGCTAGGCTC	AGCTCGAGGT	TTGTCTTACC	3840
TACATGATCA CTGCGATCCG AAGATCATTC	ACCGTGACGT	AAAAGCAGCA	AACATCCTCT	3900
TAGACGAAGA ATTOGAAGOG GTTGTTGGAG	ATTTCGGGTT	GGCAAAGCTA	ATGGACTATA	3960
AAGACACTCA CGTGACAACA GCAGTCCGTG	GCACCATCGG	TCACATCGCT	CCAGAATATC	4020
TCTCAACCGG AAAATCTTCA GAGAAAACCG	ACGITITICGG	ATACGGAATC	ATGCTTCTAG	4080
A				4081

- (2) INFORMATION FOR SEQ ID NO: 21:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 494 amino acids

(B) TYPE: amino acid

(C) STRANDEDNESS: unknown

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: protein
- (iii) HYPOTHETICAL: NO
 - (v) FRAGMENT TYPE: N-terminal
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

Met Glu Ser Ser Tyr Val Val Phe Ile Leu Leu Ser Leu Ile Leu Leu 1 5 10 15

Pro Asn His Ser Leu Trp Leu Ala Ser Ala Asn Leu Glu Gly Asp Ala 20 25 30

Leu His Thr Leu Arg Val Thr Leu Val Asp Pro Asn Asn Val Leu Gln
35 40 45

Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val Thr 50 55 60

Cys Asn Asn Glu Asn Ser Val Ile Arg Val Asp Leu Gly Asn Ala Glu 65 70 75 80

Leu Ser Gly His Leu Val Pro Glu Leu Gly Val Leu Lys Asn Leu Gln 85 90 95 WO 97/43427 PCT/EP97/02443

Glu Leu Tyr Ser Asn Asn Ile Thr Gly Pro Ile Pro Ser Asn Leu Gly 105 110 100 Asn Leu Thr Asn Leu Val Ser Leu Asp Leu Tyr Leu Asn Ser Phe Ser 125 120 115 Gly Pro Ile Pro Glu Ser Leu Gly Lys Leu Ser Lys Leu Arg Phe Leu 135 130 Arg Leu Asn Asn Asn Ser Leu Thr Gly Ser Ile Pro Met Ser Leu Thr 155 160 145 150

Asn Ile Thr Thr Leu Gln Val Leu Asp Leu Ser Asn Asn Arg Leu Ser

165 170 175

Gly Ser Val Pro Asp Asn Gly Ser Phe Ser Leu Phe Thr Pro Ile Ser 180 185 190

Phe Ala Asn Asn Leu Asp Leu Cys Gly Pro Val Thr Ser His Pro Cys
195 - 200 205

Pro Gly Ser Pro Pro Phe Ser Pro Pro Pro Pro Phe Ile Gln Pro Pro 210 225 220

Pro Val Ser Thr Pro Ser Gly Tyr Gly Ile Thr Gly Ala Ile Ala Gly
225 230 235 240

Gly Val Ala Ala Gly Ala Ala Leu Leu Phe Ala Ala Pro Ala Ile Ala 245 250 255

Phe Ala Trp Trp Arg Arg Lys Pro Leu Asp Ile Phe Phe Asp Val 260 265 270

Pro Ala Glu Glu Asp Pro Glu Val His Leu Gly Gln Leu Lys Arg Phe 275 280 285

Ser Leu Arg Glu Leu Gln Val Ala Ser Asp Gly Phe Ser Asn Lys Asn

97/43427 PCT/EP97/02443

	290					295					300				
Ile 305	Leu	Gly	Arg	Gly	Gly 310	Phe	Gly	Lys	Val	Tyr 315	Lys	Gly	Arg	Leu	Ala 320
Asp	Gly	Thr	Leu	Val 325	Ala	Val	Lys	Arg	Leu 330	Lys	Glu	Glu	Arg	Thr 335	Pro
Gly	Gly	Glu	Leu 340	Gln	Phe	Gln	Thr	Glu 345	Val	Glu	Met	Ile	Ser 350	Met	Ala
Val	His	Ar g 355	Asn	Leu	Leu	Arg	Leu 360	Arg	Gly	Phe	Cys	Met 365	Thr	Pro	Thr
Glu	A rg 370	Leu	Leu	Val	Tyr	Pro 375	Tyr	Met	Ala	Asn	Gly 380	Ser	Val	Ala	Ser
Cys 385	Leu	Arg	Glu	Arg	Pro 390	Pro	Ser	Gln	Pro	Pro 395	Leu	Asp	Trp	Pro	Thr 400
Arg	Lys	Arg	Ile	Ala 405	Leu	Gly	Ser	Ala	Arg 410	Gly	Leu	Ser	Tyr	Leu 415	His
Asp	His	Cys	Asp 420	Pro	Lys	Ile	Ile	His 425	Arg	Asp	Val	Lys	Ala 430	Ala	Asn
Ile	Leu	Leu 435	Asp	Glu	Glu	Phe	Glu 440	Ala	Val	Val	Gly	Asp 445	Phe	Gly	Leu
Ala	Lys 450	Leu	Met	Asp	Tyr	Lys 4 55	Asp	Thr	His	Val	Thr 460	Thr	Ala	Val	Arg
Gly 465	Thr	Ile	Gly	His	Ile 470	Ala	Pro	Glu	Tyr	Leu 475	Ser	Thr	Gly	Lys	Ser 480
Ser	Glu	Lys	Thr	Asp 485	Val	Phe	Gly	Tyr	Gly 490	Ile	Met	Leu	Leu		

(2)	INFORMATION	FOR	SEQ	ID	NO:	22:
-----	-------------	-----	-----	----	-----	-----

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1106 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA to mRNA

(iii) HYPOTHETICAL: NO

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 142..795

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

TOGACCCACG CGTCCGTCCA ACTTCAATAA AGGGGAAACC AACGTAACCC TAATTTTGCT 60 THETOCICIT TOTTCAGAAA ATTTTCCCTT TACTCICAAA TTCCTTTTCG ATTTCCCTCT 120 171 CTTAAACCTC CGAAAGCTCA C ATG GCG TCT CGA AAC TAT CGG TGG GAG CTC Met Ala Ser Arg Asn Tyr Arg Trp Glu Leu 5 10 TTC GCA GCT TCG TTA ACC CTA ACC TTA GCT TTG ATT CAC CTG GTC GAA 219 Phe Ala Ala Ser Leu Thr Leu Thr Leu Ala Leu Ile His Leu Val Glu 15 20 GCA AAC TCC GAA GGA GAT GCT CTC TAC GCT CTT CGC CGG AGT TTG ACA 267 Ala Asn Ser Glu Gly Asp Ala Leu Tyr Ala Leu Arg Arg Ser Leu Thr 35 40 30

																т сст	
Asī) Pr	0	Asp 45		s Va	l Le	u Gli) Ası	p Pro) Th			l As	n Pro	
			7.	,				5	U				59	5			
TGI	· AC	C	TGG	TI	C CA	r Gr	C ACC	TG	r aac	CAZ	A GAC	AA C	c cc	GIV	CAC	rcgr	363
																r Arg	
	6	0					65	5				7	0				
त्याद	GA	Ţ	באויי	GGZ	ממ ב	יראוזי יו	1 22C	. ~~~	, m-	. ~~		. ~					
																CTT Leu	411
75				-	,	80				U.J	85		, vr	PIC) GI	90	
																30	
																ATC	459
Gly	Ly	5]	Leu	Glu			Gln	Тут	Leu	Glu	Leu	Тух	Lys	Ast	Asr.	lle	
					95	5				100					105	i	
CAA	GGZ	A 2	CT	ATA	. cc1	· 100	GAA	CII	GGA	ААТ	تىلت	AAC	ያ አልጥ	حالات	יצויה י	300	507
							Glu										50 7
				110					115			-		120			
							AAT										555
Deu	W2F		.25	ığı	Asn	Asn	Asn		Thr	Gly	Ile	Val		Thr	Phe	Leu	
								130					135				
GGA	AAA	r	TG .	AAG	TCT	CTG	GTC	TTT	TTA	CGG	CTT	AAT	GAC	AAC	CGA	TTG	603
							Val										
	140						145					150					
ACC	GGT	· C	CA.	יידב	מידי	GNC	CAC	ш-л	~~	<i>C</i>	maa						
							CAC His										651
155						160		-	·Ly	9211	165	GIII	ALC.	Pile	rys	170	
																170	
GIT																	699
Val .	Asp	V	al :	Ser		Asn	Asp	Leu	Cys	Gly	Thr	Ile	Pro	Thr	Asn	Gly	
					175					180					185		
ccc '	TTT	G	er c	CAC	ATT	CCT	TTA	CAG	AAC	TTT	GAG .	AAC	AAC	CCG	AGA	TTG	747

Pro Phe Ala His Ile Pro Leu Gln Asn Phe Glu Asn Asn Pro Arg Leu	
190 195 200	
	795
GAG GGA CCG GAA TTA CTC GGT CTT GCA AGC TAC GAC ACT AAC TGC ACC	793
Glu Gly Pro Glu Leu Leu Gly Leu Ala Ser Tyr Asp Thr Asn Cys Thr	
205 210 215	
TGAAACAACT GGCAAAACCT GAAAATGAAG AATTGGGGGG TGACCTTGTA AGAACACTTC	855
ACCACTITAT CAAATATCAC ATCTATTATG TAATAAGTAT ATATATGTAG TAAAAACAAA	915
·	
AAAAATGAAG AATCGAATCG GTAATATCAT CTGGTCTCAA TTGAGAACTT CGAGGTCTGT	975
ATGIAAAATT TCTAAATGCG ATTTTCGCTT ACTGTAATGT TCGGTTGTGG GATTCTGAGA	1035
AGTAACATIT GTATIGGTAT GGTATCAAGT TGITCTGCCT TGICTGCAAA AAAAAAAAAA	1095
United prints and an annual second	
AAAAAAAAA A	1106
ACCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	

(2) INFORMATION FOR SEQ ID NO: 23:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 218 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:

Met Ala Ser Arg Asn Tyr Arg Trp Glu Leu Phe Ala Ala Ser Leu Thr 1 5 10 15

Leu Thr Leu Ala Leu Ile His Leu Val Glu Ala Asn Ser Glu Gly Asp 20 25 30

- Ala Leu Tyr Ala Leu Arg Arg Ser Leu Thr Asp Pro Asp His Val Leu 35 40 45
- Gln Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val
 50 55 60
- Thr Cys Asn Gln Asp Asn Arg Val Thr Arg Val Asp Leu Gly Asn Ser 65 70 75 80
- Asn Leu Ser Gly His Leu Ala Pro Glu Leu Gly Lys Leu Glu His Leu

 85

 90

 95
- Gln Tyr Leu Glu Leu Tyr Lys Asn Asn Ile Gln Gly Thr Ile Pro Ser 100 105 110
- Glu Leu Gly Asn Leu Lys Asn Leu Ile Ser Leu Asp Leu Tyr Asn Asn 115 120 125
- Asn Leu Thr Gly Ile Val Pro Thr Phe Leu Gly Lys Leu Lys Ser Leu 130 135 140 .
- His Ser Arg Gln Ser Gln Ala Phe Lys Val Val Asp Val Ser Ser Asn 165 170 175
- Asp Leu Cys Gly Thr Ile Pro Thr Asn Gly Pro Phe Ala His Ile Pro 180 185 190
- Leu Gln Asn Phe Glu Asn Asn Pro Arg Leu Glu Gly Pro Glu Leu Leu 195 200 205
- Gly Leu Ala Ser Tyr Asp Thr Asn Cys Thr 210 215
- (2) INFORMATION FOR SEQ ID NO: 24:

(i) SEQUENCE CHARACTERISTICS:

		(⊉) LE	NGTH	: 98	1 ba	se p	airs	3								
		(E	3) TY	PE:	nucl	eic	acid	ì									
		(0	:) SI	RANI	EINE	SS:	sing	le									
		(I) TC	POLO	GY:	line	ar										
	(ii)	MOI	ECUI	E TY	TPE:	c DN A	to.	mRNZ	A.								
	(iii)	HYI	OTHE	TIC	L: N	10											
	(ix)	FEA	ATURE	E:													
			A) NP														
		(E	3) LC	CATI	ON:	104.	.757	,									
										_							
	(xi)	SEC	DVEV	TE DE	SCRI	PTIC	2N: S	EQ 1	ID N	D: 24	1:						

AGIY	TGA C	TA A	ATTT	GTT	ල ෆ	MACI	rccro	TT	GTT	CAGA	AAA	rrr	ccc i	MTTA	MCIC	A 60	,
												ATG	ccc	TCT	CGA		
												ATG Met	ccc		CGA		
												ATG	ccc	TCT	CGA		
AAT	rccri	rr c	XGAT'I	TCCC	er ci	PCTT#	AAACC	TO	CGAA/	AGCT	CAC	ATG Met 1	GCG Ala	TCT Ser	CGA Arg	115	i
AAT.	TAT	CGG	CGATT TGG	TCCC	cic	TIC	JAACO	cct	TCG	AGCT TTA	CAC	ATG Met 1 CTA	GCG Ala ACC	TCT Ser	CGA Arg	115	i
AAT AAC Asn	TAT	CGG	CGATT TGG	TCCC	er er ere Leu	TIC	JAACO	cct	TCG	AGCT TTA Leu	CAC	ATG Met 1	GCG Ala ACC	TCT Ser	CGA Arg	115	i
AAT.	TAT	CGG	CGATT TGG	TCCC	cic	TIC	JAACO	cct	TCG	AGCT TTA	CAC	ATG Met 1 CTA	GCG Ala ACC	TCT Ser	CGA Arg GCT Ala	115	i
AAC Asn 5	TAT TYT	CGG Arg	TGG Trp	GAG Glu	CTC Leu 10	TTC Phe	GCA Ala	GCT Ala	TCG Ser	TTA Leu 15	CAC ACC Thr	ATG Met 1 CTA Leu	GCG Ala ACC Thr	TCT Ser TTA Leu	CGA Arg GCT Ala 20	115	
AAC Asn 5	TAT TYT	CAC	TGG Trp	GAG Glu GTC	CTC Leu 10 GAA	TTC Phe GCA	GCA Ala	GCT Ala	TCG Ser GAA	TTA Leu 15	CAC ACC Thr	ATG Met 1 CTA Leu	GCG Ala ACC Thr	TCT Ser TTA Leu	CGA Arg GCT Ala 20 GCT	115 163	
AAC Asn 5	TAT TYT	CAC	TGG Trp	GAG Glu GTC	CTC Leu 10 GAA	TTC Phe GCA	GCA Ala	GCT Ala	TCG Ser GAA	TTA Leu 15	CAC ACC Thr	ATG Met 1 CTA Leu	GCG Ala ACC Thr	TCT Ser TTA Leu	CGA Arg GCT Ala 20 GCT	115 163	
AAC Asn 5	TAT TYT	CAC	TGG Trp	GAG Glu GTC Val	CTC Leu 10 GAA	TTC Phe GCA	GCA Ala	GCT Ala	TCG Ser GAA Glu	TTA Leu 15	CAC ACC Thr	ATG Met 1 CTA Leu	GCG Ala ACC Thr	TCT Ser TTA Leu TAC Tyr	CGA Arg GCT Ala 20 GCT	115 163	
AAC Asn 5	TAT Tyr ATT Ile	CGG Arg CAC His	TGG Trp CTG Leu	GAG Glu GTC Val 25	CTC Leu 10 GAA Glu	TTC Phe GCA Ala	GCA Ala AAC Asn	GCT Ala TCC Ser	TCG Ser GAA Glu 30	TTA Leu 15 GGA Gly	ACC Thr GAT Asp	ATG Met 1 CTA Leu	GCG Ala ACC Thr CTC Leu	TCT Ser TTA Leu TAC Tyr 35	CGA Arg GCT Ala 20 GCT Ala	115 163	
AAC Asn 5 TTG Leu	TAT Tyr ATT Ile	CGG Arg CAC His	TGG Trp CTG Leu	GAG Glu GTC Val 25	CTC Leu 10 GAA Glu	TTC Phe GCA Ala	GCA Ala AAC Asn	GCT Ala TCC Ser	TCG Ser GAA Glu 30	TTA Leu 15 GGA Gly	CAC ACC Thr GAT Asp	ATG Met 1 CTA Leu GCT Ala	GCG Ala ACC Thr CTC Leu	TCT Ser TTA Leu TAC Tyr 35	CGA Arg GCT Ala 20 GCT Ala	115 163 211	

CCA	ACI	CIT	gri	' AAT	· cci	TGT	ACC	TGG	TIC	CAT	GIC	: ACC	TGI	' AAC	CAA	307
Pro	Thr			Asn	Pro	Cys	Thr	Trp	Phe	His	Val	Thr	Cys	Asn	Gln	
		55	,				60					65				
GAC	አአር	. ~~	. ~~~	- h (**)	~	~~~	G) M		~							
															GGA Gly	355
	70		-	••••	.49	75	qen	Deu	GIY	ASII	. Ser 80		reu	ser	GIĀ	
CAT	CIT	GCG	CCT	GAG	CTT	GGG	AAG	CIT	GAA	CAT	TTA	CAG	TAT	CTA	GAG	403
His	Leu	Ala	Pro	Glu	Leu	Gly	Lys	Leu	Glu	His	Leu	Gln	Tyr	Leu	Glu	
85					90					95					100	
						CAA										451
Leu	ŢŢŢ	Lys	Asn		Ile	Gln	Gly	Thr		Pro	Ser	Glu	Leu	Gly	Asn	
				105					110					115		
crc	AAG	AAT	CTC	ATC	AGC	TTG	GAT	CTG	TAC	AAC	AAC	ልልጥ	بلعلت	ACA	ccc	499
						Leu										477
			120				_	125	_				130		3	
ATA	GIT	ccc	ACT	TCT	TIG	GGA	AAA	TTG	AAG	TCT	CTG	GTC	TTT	TTA	CCC	547
Ile	Val		Thr	Ser	Leu	Gly	Lys	Leu	Lys	Ser	Leu	Val	Phe	Leu	Arg	
		135					140					145				
بليلت	דעע	GAC	ልልሮ	CCD.	حكلفك	ACC	~~m	~~	8.000	~~	202					
						Thr										595
	150			9		155	Gry	110	116	FLO	160	Aia	ren	ти	Ala	
ATC	CCA	AGC	CIT	AAA	GIT	GTT	GAC	GTC	TCA	AGC	AAT	GAT	TIG	TGT	GGA	643
Ile	Pro	Ser	Leu	Lys	Val	Val	Asp	Val	Ser	Ser	Asn	Asp	Leu	Cys	Gly	
165					170					175					180	
						_ ccc										691
INT	тте	Pro	'Ihr		Gly	Pro	Phe	Ala		Ile	Pro	Leu	Gln		Phe	
				185					190					195		
GAG	AAC	AAÇ	ಯಾ	AGA	TTG	GAG	GGA	ccc	GAA	TTA	cīc	GGT	CTT	GCA	AGC	739

Glu Asn Asn Pro Arg Leu Glu G	ly Pro Glu Leu Leu	ı Gly Leu Ala Ser	
200	205	210	
TAC GAC ACT AAC TGC ACC TGAAA Tyr Asp Thr Asn Cys Thr 215	CAACT GGCAAAACCT G	EAAAATGAAG	787
AATTGGGGGG TGACCTTGTA AGAACAC	TTC ACCACTITAT CAA	AATATCAC ATCTATTATG	847
TAATAAGTAT ATATATGTAG TAAAAAC	AAA AAAAATGAAG AAT	ICGAATCG GTAATATCAT	907
CIGGICICAA IIGAGAACII CGAGGIC	TGT ATGTAAAATT TC	TAAATGCG ATTTTCGCCT	967
AAATTACTCA CACT		,	981
(2) INFORMATION FOR SEQ ID NO			
(i) SEQUENCE CHARACTER (A) LENGTH: 218 ami			
(B) TYPE: amino aci			
(D) TOPOLOGY: linea	ır		
(ii) MOLECULE TYPE: prote	ein		
(xi) SEQUENCE DESCRIPTION	1: SEQ ID NO: 25:		
Met Ala Ser Arg Asn Tyr Arg T	Trp Glu Leu Phe Al	a Ala Ser Leu Thr	
1 5	10	15	
Leu Thr Leu Ala Leu Ile His I	eu Val Glu Ala As	n Ser Glu Gly Asp	
20	25	30	
Ala Leu Tyr Ala Leu Arg Arg S	Ser Leu Thr Asp Pr		
35	40	45	
Gln Ser Trp Asp Pro Thr Leu \	Val Asn Pro Cys Th	r Trp Phe His Val	

50 55 60

Thr Cys Asn Gln Asp Asn Arg Val Thr Arg Val Asp Leu Gly Asn Ser
65 70 75 80

Asn Leu Ser Gly His Leu Ala Pro Glu Leu Gly Lys Leu Glu His Leu 85 90 95

Gln Tyr Leu Glu Leu Tyr Lys Asn Asn Ile Gln Gly Thr Ile Pro Ser 100 105 110

Glu Leu Gly Asn Leu Lys Asn Leu Ile Ser Leu Asp Leu Tyr Asn Asn 115 120 125

Asn Leu Thr Gly Ile Val Pro Thr Ser Leu Gly Lys Leu Lys Ser Leu 130 135 140

Val Phe Leu Arg Leu Asn Asp Asn Arg Leu Thr Gly Pro Ile Pro Arg 145 150 155 160

Ala Leu Thr Ala Ile Pro Ser Leu Lys Val Val Asp Val Ser Ser Asn 165 170 175

Asp Leu Cys Gly Thr Ile Pro Thr Asn Gly Pro Phe Ala His Ile Pro 180 185 190

Leu Gln Asn Phe Glu Asn Asn Pro Arg Leu Glu Gly Pro Glu Leu Leu 195 200 205

Gly Leu Ala Ser Tyr Asp Thr Asn Cys Thr 210 215

- (2) INFORMATION FOR SEQ ID NO: 26:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 789 base pairs

(B) TYPE: nucleic acid

286

) ST					le								
	(ii)	MOL	ECUL	E TY	PE:	c IN A	to	mRNA								
(iii)	HYF	OTHE	MICA	L: N	10										
	(xi)	FEA														
		,	() NZA													
		(E	3) LC	XCATI	.ON:	2€	61									
	(xi)	SEC	UENC	E DE	SCRI	PTIC	INI: 5	EQ I	D NO): 26	S:					
	,,,,,							_								
тс	A CC	C AC	x	ar co	CC CC	A A	AC TA	AT CC	G TO	G GA	ks ca	C T	rc GC	CA GO	T	46
Aı	g Pr	o Tr	ır Aı	g Pr	o Ar	ng As	an T	r Ar	g Ti	æ Gj	lu Le	eu Ph	ne Al	la Al	la	
	1				5				1	.0				1	L5	
		ATC														94
Ser	Leu	Ile	Leu		Leu	Ala	Leu	He	H1S	Leu	Vai	GIU	Ala	30	Ser	
				20					25					50		
GAA	GGA	GAT	GCT	CIT	TAC	GCT	CTT	CGC	CGG	AGT	TTA	ACA	GAT	ccc	GAC	142
		Asp														
			35					40					45			
CAT	GIT	CTC	CAG	AGC	TGG	GAT	CCA	ACT	CIT	GIT	AAT	CCI	TGT	ACC	TGG	190
His	Val	Leu	Gln	Ser	Trp	Asp	Pro	Thr	Leu	Val	Asn	Pro	Cys	Thr	Trp	
		50					55					60				
	CAM	GIC	»~~	بالمحالة	220	CAA	CNC	አልሮ	~~~	حمد	אכידי	رتتا	Calc:	CAT	מייני.	238
		Val	*													230
- 116	65	V CL L		-12		70			9		75	9				
	13.3					70					, ,					

GGG AAT TCA AAC CTC TCT GGA CAT CTT GCG CCT GAG CTT GGG AAG CTT

Gly Asn Ser Asn Leu Ser Gly His Leu Ala Pro Glu Leu Gly Lys Leu

80					85					90					95		
GAA																	334
Glu	His	Leu	GIII	19r 100	Leu	GIU	Leu	TYT	105	ASTI	Asn	TTE	GIII	110	THE		
				100					103					110			
ATA	CCT	TCC	GAA	CIT	GGA	AAT	CIG	AAG	AAT	CIC	ATC	AGC	TTG	GAT	CTG		382
Ile	Pro	Ser	Glu	Leu	Gly	Asn	Leu	Lys	Asn	Leu	Ile	Ser	Leu	Asp	Leu		
			115					120					125				
TAC																	430
Tyr	ASII	130	ASII	reu	III	GIÀ	135	Val	Pro	1111	Ser	140	GIĀ	Lys	Leu		
		130					133					740					
AAG	TCT	CTG	GTC	TTT	TTA	CGG	CIT	AAT	GAC	AAC	CGA	TTG	ACG	GGG	CCA		478
Lys	Ser	Leu	Val	Phe	Leu	Arg	Leu	Asn	Asp	Asn	Arg	Leu	Thr	Gly	Pro		
	145					150					155						
					ACT												526
160	Pro	Arg	ATA	ren	Thr 165	ALA.	TTE	PTO	ser	170	rys	Val	Vai	Asp	175		
100					103					110					113		
TCA	AGC	AAT	GAT	TTG	TGT	GGA	ACA	ATC	CCA	ACA	AAC	GGA	CCT	TIT	GCT		574
Ser	Ser	Asn	Asp	Leu	Cys	Gly	Thr	Ile	Pro	Thr	Asn	Gly	Pro	Phe	Ala		
				180					185					190			
				_	AAC												622
HIS	11e	PTO	195	GIN	Asn	Phe	GIU	200	ASN	Pro	Arg	Leu	205	GIĀ	Pro		
			1,,,					200					203				
GAA	TTA	crc	GGT	CTT	GCA	AGC	TAC	GAC	ACT	AAC	TGC	ACC	TGA	AAAA	ATT		671
Glu	Leu	Leu	Gly	Leu	Ala	Ser	Tyr	Asp	Thr	Asn	Cys	Thr					
		210					215					220					
GGC	AAAA	cer (BAAA	ATGA	AG A	ATIG	GGG	g TG	ACCT	IGTA	AGA.	ACAC	TTC .	ACCA	ZTTTA'	r	731
CAA	YTATA	CAC A	ATCT	ACTA'	IG T	ATA	agta'	T AT	TATA	GTAG	TCC	AAAA	AAA .	AAAA	AAAA		789

- (2) INFORMATION FOR SEQ ID NO: 27:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 220 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

Arg Pro Thr Arg Pro Arg Asn Tyr Arg Trp Glu Leu Phe Ala Ala Ser 1 5 10 15

Leu Ile Leu Thr Leu Ala Leu Ile His Leu Val Glu Ala Asn Ser Glu 20 25 30

Gly Asp Ala Leu Tyr Ala Leu Arg Arg Ser Leu Thr Asp Pro Asp His
35 40 45

Val Leu Gln Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe
50 55 60

His Val Thr Cys Asn Gln Asp Asn Arg Val Thr Arg Val Asp Leu Gly 65 70 75 80

Asn Ser Asn Leu Ser Gly His Leu Ala Pro Glu Leu Gly Lys Leu Glu 85 90 95

His Leu Gln Tyr Leu Glu Leu Tyr Lys Asn Asn Ile Gln Gly Thr Ile 100 105 110

Pro Ser Glu Leu Gly Asn Leu Lys Asn Leu Ile Ser Leu Asp Leu Tyr 115 120 125 Asn Asn Asn Leu Thr Gly Ile Val Pro Thr Ser Leu Gly Lys Leu Lys
130 135 140

Ser Leu Val Phe Leu Arg Leu Asn Asp Asn Arg Leu Thr Gly Pro Ile 145 150 155 160

Pro Arg Ala Leu Thr Ala Ile Pro Ser Leu Lys Val Val Asp Val Ser 165 170 175

Ser Asn Asp Leu Cys Gly Thr Ile Pro Thr Asn Gly Pro Phe Ala His

Ile Pro Leu Gln Asn Phe Glu Asn Asn Pro Arg Leu Glu Gly Pro Glu 195 200 205

Leu Leu Gly Leu Ala Ser Tyr Asp Thr Asn Cys Thr 210 215 220

- (2) INFORMATION FOR SEQ ID NO: 28:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 894 base pairs

(B) TYPE: nucleic acid(C) STRANDEINESS: single

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: CINA to mRNA
- (iii) HYPOTHETICAL: NO
- (ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 1..675

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

GGA	ccc	ATT	CAA	GCC	TCC	GAA	GGG	GAC	GCT	CTT	CAC	GCG	CIT	CGC	CGG	48
Gly	Pro	Ile	Gln	Ala	Ser	Glu	Gly	Asp	Ala	Leu	His	Ala	Leu	Arg	Arg	
1				5					10					15		
						TAA										96
Ser	Leu	Ser	Asp	Pro	Asp	Asn	Val	Val	Gln	Ser	Trp	Asp	Pro	Thr	Leu	
			20					25					30			
بلملت	ልልጥ	Ст.	ىلخىل	۳	mcc.	TTT	СУТ	حتلات	لما√∀	ىلخىك	ידממ	CAA	CAC	САТ	CAA	144
						Phe						_			_	
٧۵٢	7611	35	Cys			1110	40	,		-,-		45			~	
		-														
GIC	ACT	CGT	crg	GAT	TTG	GGG	AAT	TCA	AAC	TTA	TCT	GGA	CAT	CTA	GTA	192
Val	Thr	Arg	Leu	Asp	Leu	Gly	Asn	Ser	Asn	Leu	Ser	Gly	His	Leu	Val	
	50					55					60					
ccr	GAA	CTT	GGG	AAG	CTT	GAA	CAT	TTA	CAA	TAT	CIG	TAT	GGA	ATC	ATC	240
Pro	Glu	Leu	Gly	Lys	Leu	Glu	His	Leu	Gln	Tyr	Leu	Tyr	Gly	Ile	Ile	
65					70					75					80	
ACT	CIT	TTG	CCT	TTT	GAT	TAT	CTG	AAA	ACA	TIT	ACA	TTA	TCA	GIC	ACA	288
Thr	Leu	Leu	Pro	Phe	Asp	Tyr	Leu	Lys	Thr	Phe	Thr	Leu	Ser	Val	Thr	
				85					90					95		
																226
						GAG										336
HIS	iie	ınr		Cys	Pne	Glu	ser		Ser	GIU	Ten	ıyr		ASII	GIU	
			100					105					110			
ATT	CAA	GGA	ACT	ATA	CCT	TCT	GAG	CTT	GGA	AAT	CTG	AAG	AGT	CTA	ATC	384
						Ser									_	
		115					120		_			125				
agt	TTG	GAT	CTG	TAC	AAC	AAC	AAT	CTC	ACC	GGG	AAA	ATC	CCA	TCT	TCT	432
Ser	Leu	Asp	Leu	Tyr	Asn	Asn	Asn	Leu	Thr	Gly	Lys	Ile	Pro	Ser	Ser	
	130					135					140					

TTG	GGA	AAA	TTG	AAG	TCA	CIT	GTT	TTT	TTG	ccc	CTT	AAC	GAA	AAC	CGA	480
Leu	Gly	Lys	Leu	Lys	Ser	Leu	Val	Phe	Leu	Arg	Leu	Asn	Glu	Asn	Arg	
145					150					155					160	
TTG	ACC	GGT	CCT	ATT	CCT	AGA	GAA	CIC	ACA	GIT	ATT	TCA	AGC	CIT	AAA	528
Leu	Thr	Gly	Pro	Ile	Pro	Arg	Glu	Leu	Thr	Val	Ile	Ser	Ser	Leu	Lys	
				165					170					175		
CIT	GIT	GAT	GIC	TCA	GGG	AAT	GAT	TTG	TGT	GGA	ACA	ATT	CCA	GTA	GAA	576
Val	Val	Asp	Val	Ser	Gly	Asn	Asp	Leu	Cys	Gly	Thr	Ile	Pro	Val	Glu	
			180					185					190			
GGA	CCT	TTT	GAA	CAC	TTA	CCT	ATG	CAA	AAC	TTT	GAG	AAC	AAC	CTG	AGA	624
Gly	Pro	Phe	Glu	His	Ile	Pro	Met	Gln	Asn	Phe	Glu	Asn	Asn	Leu	Arg	
		195					200					205				
TTG	GAG	GGA	CCA	GAA	CTA	CTA	GGT	CTT	GCG	AGC	TAT	GAC	ACC	AAT	TGC	672
Leu	Glu	Gly	Pro	Glu	Leu	Leu	Gly	Leu	Ala	Ser	Tyr	Asp	Thr	Asn	Cys	
	210					215					220					
															•	
ACT	TAAA	AAGA	AG I	TGAA	GAAC	C TA	4AAT	GAAC	CAA :	GIT	GCT	GACC	TIG	AA		725
Thr																
225																
GAAC	TCTG	TA C	CAAG	IGII	T GI	TAAAT	CTAT	ATA	GAGO	CIT	GITI	CATO	TT A	TATA	AAADT	785
GCTT	TGAG	AG A	CAGI	'AAC'I	T GC	CAATO	TATI	GGI	ATTC	GTA	GAAA	AAGI	TG A	AATC	AGAAT	845
ICCI	TIGI	r aa	TGGA	TTTG	ar Gi	MCI	TATO	TAA	CTTC	AAT	TICI	TATI	'A			894

(2) INFORMATION FOR SEQ ID NO: 29:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 225 amino acids

(B) TYPE: amino acid(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SE	QUENCE D	ESCRIPTI	ON: SEX) ID 1/4	D: 29:			
Gly Pro Ile	Gln Ala	Ser Glu	Gly As	ap Ala 10	Leu His	s Ala	Leu Arg	
Ser Leu Ser	Asp Pro 20	Asp Asn		al Gln 25	Ser Tr	Asp	Pro Thr	Leu
Val Asn Pro	Cys Thr	Trp Phe	His Va	ıl Thr	Cys Asi	Gln 45	His His	Gln
Val Thr Arg 50	Leu Asp	Leu Gly	Asn Se	er Asn	Leu Sei		His Leu	Val
Pro Glu Leu 65	Gly Lys	Leu Glu 70	His Le	eu Gln	Tyr Lei 75	Tyr	Gly Ile	Ile 80
Thr Leu Leu	Pro Phe 85	Asp Tyr	Leu Ly	rs Thr 90	Phe Thr	: Leu	Ser Val	
His Ile Thr	Phe Cys	Phe Glu	Ser Ty	_	Glu Leu	ıTyr	Lys Asn 110	Glu
Ile Gln Gly 115	Thr Ile	Pro Ser	Glu Le	u Gly	Asn Leu	Lys 125	Ser Leu	Ile
Ser Leu Asp	Leu Tyr	Asn Asn 135	Asn Le	u Thr	Gly Lys		Pro Ser	Ser

150

165

Leu Gly Lys Leu Lys Ser Leu Val Phe Leu Arg Leu Asn Glu Asn Arg

Leu Thr Gly Pro Ile Pro Arg Glu Leu Thr Val Ile Ser Ser Leu Lys

160

175

155

170

Val Val Asp Val Ser Gly Asm Asp Leu Cys Gly Thr Ile Pro Val Glu 180 185 190

Gly Pro Phe Glu His Ile Pro Met Gln Asn Phe Glu Asn Asn Leu Arg 195 200 205

Leu Glu Gly Pro Glu Leu Leu Gly Leu Ala Ser Tyr Asp Thr Asn Cys 210 215 220

Thr

225

- (2) INFORMATION FOR SEQ ID NO: 30:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1063 base pairs

(B) TYPE: nucleic acid(C) STRANDETNESS: single

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: CDNA to mRNA
- (iii) HYPOTHETICAL: NO
- (ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 106..759

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

TOGACCCACG CGICCGACGA AACCCTAATT TIGCTTCCTC ATCTTGTTCA GAAAATTACT

CAAATTCCTA TTAGATTACT CTCTCTTCGA CCTCCGATAG CTCAC ATG GCG TCT 114

Met Ala Ser

						CTC											162
Arg	Asn	Tyr	Arg	Trp	Glu	Leu	Phe	Ala	Ala	Ser	Leu	Ile	Leu	Thr	Leu		
	5					10					15						
GCT	TTG	ATT	CAC	CIG	arc	GAA	GCA	AAC	TCC	GAA	GGA	GAT	GCT	CIT	TAC		210
Ala	Leu	Ile	His	Leu	Val	Glu	Ala	Asn	Ser	Glu	Gly	Asp	Ala	Leu			
20					25					30					35		
						ACA											258
Ala	Leu	Arg	Arg	Ser	Leu	Thr	Asp	Pro		His	Val	Leu	Gln		Trp		
				40					45					50			
						CCT											306
Asp	Pro	Thr		Val	Asn	Pro	Cys		Trp	Phe	His	Val		Cys	Asņ		-
			55					60					65				
						CGT											354
Gln	Asp		Arg	Val	Thr	Arg		Asp	Leu	Gly	Asn		Asn	Leu	Ser		
		70					75					80					
																	400
						CIT											402
Gly		Leu	Ala	Pro	Glu	Leu	Gly	Lys	Leu	Glu		Leu	GIn	Tyr	Leu		
	85					90					95						
														~~~	001		450
						ATC									_	•	450
	Leu	Tyr	Lys	Asn		Ile	GIn	GIY	Thr		Pro	Ser	GIU	Leu			
100					105					110					115		
	~~~			~~~				~> ~	~	<b>~</b>				~			400
						AGC									_		498
ASn	Leu	rys	ASII		TIE	Ser	Leu	ASD		тут	ASII	ASII	ASI		1111		
				120					125					130			
~~·	አመጻ	بنعتم	~~	لمف لا	بلحك	TTG	CC X	አአአ	-ريس	244	بلمك	سب	ميت	بلعلف	מידינוי		546
							_						_				740
αīλ	TTE	vai		m	ser	Leu	GΤĀ	_	ren	гÃ2	ser.	ren		LINE	Den		
			135					140					145				

CGG	CIT	AAT	GAC	AAC	CGA	TIG	ACG	GGG	CCA	ATC	CCI	AGA	GCA	CIC	ACT	594
Arg	Leu	Asn	Asp	Asn	Arg	Leu	Thr	Gly	Pro	Ile	Pro	Arg	Ala	Leu	Thr	
		150					155					160				
GCA	ATC	CCA	AGC	CTT	AAA	GIT	GTT	GAT	GTC	TCA	AGC	AAT	GAT	TIG	TGT	642
Ala	Ile	Pro	Ser	Leu	Lys	Val	Val	Asp	Val	Ser	Ser	Asn	Asp	Leu	Cys	
	165					170					175					
GGA	ACA	ATC	CCA	ACA	AAC	GGA	CCT	TTT	GCT	CAC	ATT	CCT	TTA	CAG	AAC	690
Gly	Thr	Ile	Pro	Thr	Asn	Gly	Pro	Phe	Ala	His	Ile	Pro	Leu	Gln	Asn	
180					185					190					195	
TTT	GAG	AAC	AAC	∞	AGG	TTG	GAG	GGA	α	GAA	TTA	CTC	GGT	CTT	GCA	738
Phe	Glu	Asn	Asn	Pro	Arg	Leu	Glu	Gly	Pro	Glu	Leu	Leu	Gly	Leu	Ala	
				200					205					210		
AGC	TAC	GAC	ACT	AAC	TGC	ACC	TGAA	AAAA	TT G	GCAA	AACC	T GA	LAAAT	CAAC	;	789
Ser	Tyr	Asp	Thr	Asn	Cys	Thr										
			215													
AATT	GGGG	GG T	GACC	TIGI	'A AG	AACA	CTTC	: ACC	ACTI	TAT	CAAA	TATC	AC A	ATCTA	CTATG	849
TAAT	AAGT	A TA	TATA	TGTA	G TC	CAAA	AAAA	AAA	TGAA	GAA	TCGA	ATCA	CT P	IATA!	CATCI	909
GGIC	TCAA	TT G	AGAA	CITI	G AG	GICI	GIGI	' ATG	TAAA	ATT	TCTA	AATG	CG A	CITI	CCCI	969
· ~~~																
ACIG	TAAT	GT T	CGGI	IGIG	G GA	TTCT	GAGA	AGI	AACA	TTT	GTAT	TGGI	AT G	GTAT	CAAGT	1029
18 4111		~ ~							_							
1611	ديور	CT T	GICI	GCAA	A AA	AAAA	AAAA	AAA	A							1063

(2) INFORMATION FOR SEQ ID NO: 31:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 218 amino acids

(B) TYPE: amino acid

(D)	TOPOLOGY:	linear
-----	-----------	--------

(ii	.)	MOL	ECULE	TYPE:	pro	tein
-----	----	-----	--------------	-------	-----	------

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:

Met Ala Ser Arg Asn Tyr Arg Trp Glu Leu Phe Ala Ala Ser Leu Ile 1 5 10 15

Leu Thr Leu Ala Leu Ile His Leu Val Glu Ala Asn Ser Glu Gly Asp 20 25 30

Ala Leu Tyr Ala Leu Arg Arg Ser Leu Thr Asp Pro Asp His Val Leu 35 40 45

Gln Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val
50 55 60

Thr Cys Asn Gln Asp Asn Arg Val Thr Arg Val Asp Leu Gly Asn Ser 65 70 . 75 80

Asn Leu Ser Gly His Leu Ala Pro Glu Leu Gly Lys Leu Glu His Leu 85 90 95

Gln Tyr Leu Glu Leu Tyr Lys Asn Asn Ile Gln Gly Thr Ile Pro Ser 100 105 110

Glu Leu Gly Asn Leu Lys Asn Leu Ile Ser Leu Asp Leu Tyr Asn Asn 115 120 125

Asn Leu Thr Gly Ile Val Pro Thr Ser Leu Gly Lys Leu Lys Ser Leu 130 135 140

Val Phe Leu Arg Leu Asn Asp Asn Arg Leu Thr Gly Pro Ile Pro Arg 145 150 155 160

Ala Leu Thr Ala Ile Pro Ser Leu Lys Val Val Asp Val Ser Ser Asn

- 90 -

165 170 175

Asp Leu Cys Gly Thr Ile Pro Thr Asn Gly Pro Phe Ala His Ile Pro 180 185 190

Leu Gln Asn Phe Glu Asn Asn Pro Arg Leu Glu Gly Pro Glu Leu Leu 195 200 205

Gly Leu Ala Ser Tyr Asp Thr Asn Cys Thr 210 215

- (2) INFORMATION FOR SEQ ID NO: 32:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2089 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA to mRNA
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Arabidopsis thaliana
 - (vii) IMMEDIATE SOURCE:
 - (B) CLONE: SERK gene cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 195..2069

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 32:

GGATTITTAT TITATTITT ACTOPTIGIT TGTTTAATG CTAATGGGT TTTAAAAGGG	60
TTATCGAAAA AATGAGTGAG TTTGTGTTGA GGTTGTCTCT GTAAAGTGTT AATGGTGGTG	120
ATTITICGGAA GITAGGGITT TCTCGGATCT GAAGAGATCA AATCAAGATT CGAAATITAC	180
CATTGITGIT TGAA ATG GAG TCG AGT TAT GTG GTG TTT ATC TTA CTT TCA Met Glu Ser Ser Tyr Val Val Phe Ile Leu Leu Ser 1 5 10 '	230
CTG ATC TTA CTT CCG AAT CAT TCA CTG TGG CTT GCT TCT GCT AAT TTG Leu Ile Leu Leu Pro Asn His Ser Leu Trp Leu Ala Ser Ala Asn Leu 15 20 25	278
GAA GGT GAT GCT TTG CAT ACT TTG AGG GTT ACT CTA GTT GAT CCA AAC Glu Gly Asp Ala Leu His Thr Leu Arg Val Thr Leu Val Asp Pro Asn 30 35 40	326
AAT GTC TTG CAG AGC TGG GAT CCT ACG CTA GTG AAT CCT TGC ACA TGG Asn Val Leu Gln Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp 45 50 55 60	374
TTC CAT GTC ACT TGC AAC AAC GAG AAC AGT GTC ATA AGA GTT GAT TTG Phe His Val Thr Cys Asn Asn Glu Asn Ser Val Ile Arg Val Asp Leu 65 70 75	422
GGG AAT GCA GAG TTA TCT GGC CAT TTA GTT CCA GAG CTT GGT GTG CTC Gly Asn Ala Glu Leu Ser Gly His Leu Val Pro Glu Leu Gly Val Leu 80 85 90	470
AAG AAT TIG CAG TAT TIG GAG CIT TAC AGT AAC AAC ATA ACT GGC CCG Lys Asn Leu Gln Tyr Leu Glu Leu Tyr Ser Asn Asn Ile Thr Gly Pro 95 100 105	518
ATT CCT AGT AAT CTT GGA AAT CTG ACA AAC TTA GTG AGT TTG GAT CTT	566

Ile	Pro	Ser	Asn	Leu	Gly	Asn	Leu	Thr	Asn	Leu	Val	Ser	Leu	Asp	Leu	
	110					115					120					
TAC	TTA	AAC	AGC	TTC	TCC	GGT	CCT	ATT	CCG	GAA	TCA	TTG	GGA	AAG	CIT	614
Tyr	Leu	Asn	Ser	Phe	Ser	Gly	Pro	Ile	Pro	Glu	Ser	Leu	Gly	Lys	Leu	
125					130					135					140	
TCA	AAG	CTG	AGA	TIT	CTC	CGG	CTT	AAC	AAC	AAC	AGT	crc	ACT	GGG	TCA	662
Ser	Lys	Leu	Arg	Phe	Leu	Arg	Leu	Asn	Asn	Asn	Ser	Leu	Thr	Gly	Ser	
				145					150					155		
															•	
ATT	CCT	ATG	TCA	CTG	ACC	AAT	ATT	ACT	ACC	CTT	CAA	GTG	TTA	GAT	CTA	710
Ile	Pro	Met	Ser	Leu	Thr	Asn	Ile	Thr	Thr	Leu	Gln	Val	Leu	Asp	Leu	
			160					165					170			
TCA	AAT	AAC	AGA	CTC	TCT	GGT	TCA	GIT	CCT	GAC	AAT	GGC	TCC	TTC	TCA	758
Ser	Asn	Asn	Arg	Leu	Ser	Gly	Ser	Val	Pro	Asp	Asn	Gly	Ser	Phe	Ser	
		175					180					185				
						TTT										806
Leu	Phe	Thr	Pro	Ile	Ser	Phe	Ala	Asn	Asn	Leu	Asp	Leu	Cys	Gly	Pro	
	190					195					200					
						CCI										854
	Thr	Ser	His	Pro	Cys	Pro	Gly	Ser	Pro	Pro	Phe	Ser	Pro	Pro	Pro	
205					210					215					220	
						CCA							TAT	GGT	ATA	902
Pro	Phe	Ile			Pro	Pro	Val	Ser	Thr	Pro	Ser	Gly	Tyr	Gly	Ile	
				225					230					235		
						GGA										950
ınr	GIĀ	ALA		Ala	Gly	Gly			Ala	Gly	Ala	Ala	Leu	Pro	Phe	
			240					245					250			
	~~	~	~~-													
						TTT Phe										998

:	255	260)				265			
GAT ATT 1										1046
GGA CAG (Gly Gln 1285										1094
GGG TTT A			Gly							1142
TAC AAG O	_					_				1190
AAG GAA 0 Lys Glu 0			Glu							1238
GAG ATG A Glu Met I 350										1286
TTC TGT A Phe Cys M 365				Leu						1334
AAT GGA A			Arg	GAG .	AGG				ccr	1382
CCG CTT G								GCT		1430

GGT	TTG	TCT	TAC	CTA	CAT	GAT	CAC	TGC	GAT	ccc	AAG	ATC	ATT	CAC	CCI	1478
Gly	Leu	Ser	Tyr	Leu	His	Asp	His	Cys	Asp	Pro	Lys	Ile	Il	His	Arg	
		415					420					425				
GAC	GTA	AAA	GCA	GCA	AAC	ATC	crc	TTA	GAC	GAA	GAA	TIC	GAA	GCG	GIT	1526
Asp	Val	Lys	Ala	Ala	Asn	Ile	Leu	Leu	Asp	Glu	Glu	Phe	Glu	Ala	Val	
	430					435					440					
GIT	GGA	GAT	TTC	GGG	TTG	GCA	AAG	CII	ATG	GAC	TAT	AAA	GAC	ACT	CAC	1574
Val	Gly	Asp	Phe	Gly	Leu	Ala	Lys	Leu	Met	Asp	Tyr	Lys	Asp	Thr	His	
445					450					455					460	
GIG	ACA	ACA	GCA	GTC	CGT	GGC	ACC	ATC	GGT	CAC	ATC	GCT	CCA	GAA	TAT	1622
Val	Thr	Thr	Ala	Val	Arg	Gly	Thr	Ile	Gly	His	Ile	Ala	Pro	Glu	Tyr	
				465					470					475		
crc	TCA	ACC	GGA	AAA	TCT	TCA	GAG	AAA	ACC	GAC	GIT	TTC	GGA	TAC	GGA	1670
						Ser										
			480					485		_			490	•		
ATC	ATG	CTT	CTA	GAA	CTA	ATC	ACA	GGA	CAA	AGA	GCT	TTC	GAT	CIC	GCT	1718
Ile	Met	Leu	Leu	Glu	Leu	Ile	Thr	Gly	Gln	Arg	Ala	Phe	Asp	Leu	Ala	
		495					500					505				
CGG	CTA	GCT	AAC	GAC	GAC	GAC	GIC	ATG	TTA	CTT	GAC	TGG	GTG	AAA	GGA	1766
Arg	Leu	Ala	Asn	Asp	Asp	Asp	Val	Met	Leu	Leu	Asp	Trp	Val	Lys	Gly	
	510					515					520			_	-	
TIG	TTG	AAG	GAG	AAG	AAG	CTA	GAG	ATG	TTA	GTG	GAT	CCA	GAT	CTT	CAA	1814
Leu																
525					530					535	- · E -				540	
ACA	AAC	TAC	GAG	GAG	AGA	GAA	CIG	GAA	CAA	GIG	ATA	CAA	GIG	CCC	TTG	1862
Thr	Asn	Tyr	Glu	Glu	Arg	Glu	Leu	Glu	Gln	Val	Ile	Gln	Val	Ala	Leu	
				545					550					555		

CTA	TGC	ACG	CAA	GGA	TCA	CCA	ATG	GAA	AGA	CCA	AAG	ATG	TCT	GAA	GTT	1910
Leu	Cys	Thr	Gln	Gly	Ser	Pro	Met	Glu	Arg	Pro	Lys	Met	Ser	Glu	Val	
			560					565					570			
GTA	AGG	ATG	CTG	GAA	GGA	GAT	GGG	CIT	GCG	GAG	AAA	TGG	GAC	GAA	TGG	1958
Val	Arg	Met	Leu	Glu	Gly	Asp	Gly	Leu	Ala	Glu	Lys	Trp	Asp	Glu	Trp	
		575					580					585				
CAA	AAA	GIT	GAG	ATT	TIG	AGG	GAA	GAG	ATT	GAT	TIG	AGT	CCT	AAT	CCT	2006
Gln	Lys	Val	Glu	Ile	Leu	Arg	Glu	Glu	Ile	Asp	Leu	Ser	Pro	Asn	Pro	
	590					595					600					
AAC	TCT	GAT	TGG	ATT	CTT	GAT	TCT	ACT	TAC	AAT	TTG	CAC	GCC	GIT	GAG	2054
Asn	Ser	Asp	Trp	Ile	Leu	Asp	Ser	Thr	Tyr	Asn	Leu	His	Ala	Val	Glu	
605					610					615					620	
TTA	TCT	GGT	CCA	AGG	TAAA	AAAA	AA A	AAA	AAA	A						2089
Leu	Ser	Gly	Pro	Arg												
				625												

- (2) INFORMATION FOR SEQ ID NO: 33:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 625 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33:

Met Glu Ser Ser Tyr Val Val Phe Ile Leu Leu Ser Leu Ile Leu Leu 1 5 10 15

Pro Asn His Ser Leu Trp Leu Ala Ser Ala Asn Leu Glu Gly Asp Ala 20 25 30

Leu	His	Thr	Leu	Arg	Val	Thr	Leu	Val	Asp	Pro	Asn	Asn	Val	Leu	Gln
		35					40					45			
			•												
Ser	Trp	Asp	Pro	Thr	Leu	Val	Asn	Pro	Cys	Thr	Trp	Phe	His	Val	Thr
	50					55					60				
_	Asn	Asn	Glu	Asn		Val	Ile	Arg	Val	_	Leu	Gly	Asn	Ala	
65					70					75					80
•		a 1	*** =	•	• • • •		6 3	•	6 1				_		- 33
Leu	ser	GIĀ	HIS		Vai	Pro	GII	Leu	_	Vai	Leu	Lys	ASD		GIN
				85					90					95	
ጥም	I 🚓 1 1	Ghi	Len	Тълг	Ser	Asn	Δen	Tla	ጥኮታ	Gly	Pro	Tla	Pro	Sar	λen
-7-		014	100	-7-	J-L	71,,,11	~~~	105	****	CLy	110	110	110	ے۔	101,
Leu	Gly	Asn	Leu	Thr	Asn	Leu	Val	Ser	Leu	Asp	Leu	Tyr	Leu	Asn	Ser
		115					120			_		125			
Phe	Ser	Gly	Pro	Ile	Pro	Glu	Ser	Leu	Gly	Lys	Leu	Ser	Lys	Leu	Arg
	130					135					140				
Phe	Leu	Arg	Leu	Asn	Asn	Asn	Ser	Leu	Thr	Gly	Ser	Ile	Pro	Met	Ser
145					150					155					160
Leu	Thr	Asn	Ile	Thr	Thr	Leu	Gln	Val	Leu	Asp	Leu	Ser	Asn	Asn	Arg
				165					170					175	
	_		_		_	_			_		_				
ren	Ser	GIY		Val	Pro	Asp	Asn		Ser	Phe	Ser	Leu		Thr	Pro
			180					185					190		
Tle	Cor	Dhe	21-	١	١	1		7	^	01	D	**- 1	Man-	C	77
TTG	ser	195	ATG	ASI	ASTI	Leu	200	∟eu	cys	GIĀ	PTO	205	ınr	ser	HIS
		122					ZUU					ZUJ			

Pro Cys Pro Gly Ser Pro Pro Phe Ser Pro Pro Pro Pro Phe Ile Gln

220

215

Pro 225	Pro	Pro	Val	Ser	Thr 230	Pro	Ser	Gly	Tyr	G1y 235	IIe	Thr	GIĀ	ALA	240
Ala	Gly	Gly	Val	Ala 245	Ala	Gly	Ala	Ala	Leu 250	Pro	Phe	Ala	Ala	Pro 255	Ala
Ile	Ala	Phe	Ala 260	Trp	Trp	Arg	Arg	Arg 265	Ser	Pro	Leu	Asp	Ile 270	Phe	Phe
Asp	Val	Pro 275	Ala	Glu	Glu	Asp	Pro 280	Glu	Val	His	Leu	Gly 285	Gln	Leu	Lys
Arg	Phe 290	Ser	Leu	Arg	Glu	Leu 295	Gln	Val	Ala	Ser	Aട p 300	Gly	Phe	Ser	Asn
Lys 305	Asn	Ile	Leu	Gly	Arg 310	Gly	Gly	Phe	Gly	Lys 315	Val	Tyr	Lys	Gly	Arg 320
Leu	Ala	Asp	Gly	Thr 325	Leu	Val	Ala	Val	Lys 330	Arg	Leu	Lys	Glu	Glu 335	Arg
Thr	Pro	Gly	Gly 340	Glu	Leu	Gln	Phe	Gln 345	Thr	Glu	Val	Glu	Met 350	Ile	Ser
Met	Ala	Val 355	His	Arg	Asn	Leu	Leu 360	Arg	Leu	Arg	Gly	Phe 365	Cys	Met	Thr
Pro	Thr 370	Glu	Arg	Leu	Leu	Val 375	Tyr	Pro	Tyr	Met	Ala 380	Asn	Gly	Ser	Val
Ala 385	Ser	Cys	Leu	Arg	Glu 390	Arg	Pro	Pro	Ser	Gln 395	Pro	Pro	Leu	Asp	Trp 400
Pro	Thr	Arg	Lys	Arg 405	Ile	Ala	Leu	Gly	Ser 410	Ala	Arg	Gly	Leu	Ser 415	Tyr
Leu	His	Asp	His	Cys	Asp	Pro	Lys	Ile	Ile	His	Arg	Asp	Val	Lys	Ala

. .

- 98 -

			420					425					430		
Ala	Asn	Ile 435	Leu	Leu	Asp	Glu	Glu 440	Phe	Glu	Ala	Val	Val 445	Gly	Asp	Phe
Gly	Leu 450	Ala	Lys	Leu	Met	А s p 455	Tyr	Lys	Asp	Thr	His 460	Val	Thr	Thr	Ala
Val 465	Arg	Gly	Thr	Ile	Gly 4 70	His	Ile	Ala	Pro	Glu 475	Tyr	Leu	Ser	Thr	Gly 480
Lys	Ser	Ser	Glu	Lys 485	Thr	Asp	Val	Phe	Gly 490	Tyr	Gly	Ile	Met	Leu 495	Leu
Glu	Leu	Ile	Thr 500	Gly	Gln	Arg	Ala	Phe 505	Asp	Leu	Ala	Arg	Leu 510	Ala	Asn
Asp	Asp	Asp 515	Val	Met	Leu	Leu	Asp 520	Trp	Val	Lys	Gly	Leu 525	Leu	Lys	Glu
Lys	Lys 530	Leu	Glu	Met	Leu	Val 535	Asp	Pro	Asp	Leu	Gln 540	Thr	Asn	Tyr	Glu
Glu 545	Arg	Glu	Leu	Glu	Gln 550	Val	Ile	Gln	Val	Ala 555	Leu	Leu	Cys	Thr	Gln 560
Gly	Ser	Pro	Met	Glu 565	Arg	Pro	Lys	Met	Ser 570	Glu	Val	Val	Arg	Met 575	Leu
Glu	Gly	Asp	Gly 580	Leu	Ala	Glu	Lys	Trp 585	Asp	Glu	Trp	Gln	Lys 590	Val	Glu
Ile	Leu	Arg 595	Glu	Glu	Ile	Asp	Leu 600	Ser	Pro	Asn	Pro	Asn 605	Ser	Asp	Trp
Ile	Leu 610	Asp	Ser	Thr	Tyr	Asn 615	Leu	His	Ala	Val	Glu 620	Leu	Ser	Gly	Pro

WO 97/43427 PCT/EP97/02443

- 99 -

Arg

REFERENCES

Aleith, F. and Richter, G. (1990) Planta 183, 17-24.

Bent, A.F., Kunkel, B.N., Dahlbeck D., Brown, K.L., Schmidt, R., Giraudat, J., Leung, J. and Staskawicz, B.J. (1994) Science 265, 1856-1860.

Braun T, Schofield P.R. and Sprengel R. (1991) EMBO J. 10, 1885-1890.

Dangi, J.L. (1995) Cell 80, 363-366.

De Jong, A.J., Schmidt, E.D.L. and De Vries, S.C. (1993) Plant. Mol. Biol. 5, 367-377.

De Vries, S.C., Booij, H., Meyerink, P., Huisman, G., Wilde, H.D., Thomas T.L. and Van Kammen, A. (1988a) Planta 176, 196-204.

De Vries, S.C., Hoge, H., and Bisseling, T. (1988b) In Plant Molecular Biology Manual, B6,

S.B. Gelvin, R.A. Schilperoort, and D.P.S. Verma. eds (Dordrecht, the Netherlands: Kluwer Academic Publishers), pp 1-13.

Dubois, T., Guedira, M., Dubois, D. and Vasseur, J. (1991) Protoplasma 162, 120-127.

Dudits, D., Györgyey, J., Bögre, L. and Bakó, L. (1995) In: In vitro Embryogenesis in Plants. Ed. Thorpe, T.A. Kluwer Press, pp. 267-308.

Engler J.A., Van Montagu M. and Engler G. (1994) Plant Mol. Biol. Rep. 12, 321-331.

Gamborg, O.L., Miller, R.A. and Ojima, K. (1968) Exp. Cell. Res. 50, 151-158.

Giorgetti, L., Vergara, M.R., Evangelista, M., LoSchiavo, F., Terzi, M. and Ronchi, V.N. (1995) Mol. Gen. Genet. 246, 657-662.

Goldberg, R.B., Barker, S.J. and Perez-Grau, L. (1989) Cell 56, 149-160.

Goldberg, R.B., Barker, S.J., Perez-Grau, L. (1989) Cell 56, 149-160.

Goldberg, R.B., de Paiva, G. and Yadegari, R. (1994) Science 266, 605-614.

Govind, S. and Steward, R. (1991) Dorsoventral pattern formation in *Drosophila*. Trends Genet. 7, 119-125.

Guzzo, F., Baldan, B., Levi, M., Sparvoli, E., LoSchiavo, F., Terzi, M., and P. Mariani (1995). Protoplasma 185, 28-36.

Guzzo, F., Baldan, B., Mariani, P., LoSchiavo, F. and Terzi, M. (1994) Exper. Botany 45, 1427-1432.

Hanks, S.K., Quinn, A.M. and Hunter, T. (1988) Science 241, 42-52.

Hashimoto, C., Hudson, K.L. and Anderson K.V. (1988) Cell 52, 269-279.

Heck, G.R., Perry, S.E., Nichols, K.W. and Fernandez, D.E. (1995) AGL15, a MADS domain protein expressed in developing embryos. Plant Cell 7, 1271-1282.

Heldin, C-H. (1995) Cell 80, 213-223.

Hodge, R., Paul, Wyatt, Draper, J. and Scott, R. (1992) Plant J. 2, 257-260.

Hom, M.A. and Walker, J.C. (1994) Biochim. Biophys. Acta 1208, 65-74.

Jones, D.A., Thomas, C.M., Hammond-Kosack, K.E., Balint-Kurti, P.J. and Jones, J.D.G. (1994) Science 266, 789-792.

Kobe B. and Deisenhofer J. (1994) TIBS 19, 415-421.

Li, F., Barnathan, E.S. and Karikó, K. (1994) Nucl. Acid Res. 22, 1764-1765.

Liang, P. and Pardee, A.B. (1992) Science, 257, 967-971.

Meyerowitz, E.M. (1995) EDBC 95 congress of the European developmental biology organization. Abstract SI4.

Mu J-H., Lee H-S. and Kao T-h. (1994) Plant Cell 6, 709-721.

Pennell, R.I., Janniche, L., Scofield, G.N., Booij, H., de Vries, S.C. and Roberts, K. (1992) J. Cell Biol. 119, 1371-1380.

Rounsley, S.D., Ditta, G.S. and Yanofsky, M.F. (1995) Plant Cell 7, 1259-1269.

Sato, S., Toya, T., Kawahara, R., Whittier, R.F., Fukuda, H. and Komamine, A. (1995) Plant Mol. Biol. 28, 39-46.

Shelton, C.A. and Wasserman, S.A. (1993) Cell 72, 515-525.

Sterk, P., and De Vries, S.C. (1992) In Redenbaugh K (ed), Synseeds: Applications of synthetic seeds to crop improvement, CRC Press, London (1992).

Sterk, P., Booij, H., Schellekens, G.A., Van Kammen, A. and De Vries, S.C. (1991) Plant Cell 3, 907-921.

Thomas, T.L. (1993) Plant Cell 5, 1401-1410

Toonen, M.A.J. and De Vries, S.C. (1995) In: Embryogenesis, the generation of a plant.

Eds. Wang, T.L. and Cuming, A. BIOS Scientific Publishers, Oxford, UK, pp. 173-189.

Toonen, M.A.J., Hendriks, T., Schmidt, E.D.L., Verhoeven, H.A., Van Kammen, A and De Vries, S.C. (1994) Planta 194, 565-572.

Toonen, M.A.J., Schmidt, E.D.L., Hendriks, T., Verhoeven, H.A., Van Kammen, A. and De Vries, S.C. (1996) *submitted*.

Torii, K.U. and Komeda, Y. (1994) 4th international congress of plant molecular biology. Abstract 692,

Van Engelen, F.A. and De Vries, S.G. (1992) Trends Genet. 8, 66-70.

Vamer, J.E. and Lin, L.-S. (1989) Cell 56, 231-239.

Walker J.C. (1994) Plant Mol. Biol. 26, 1599-1609.

Walker, J.C. (1993) Plant J. 3, 451-456.

Whitham, S., Dinesh-Kumar, S.P., Choi, D., Hehl, R., Corr, C. and Baker, B. (1994) Cell 78, 1101-1115.

Wilde, H.T., Nelson, W.S., Booij H., de Vries, S.C. and Thomas, T.L. (1988) Planta 176, 205-213.

Wurtele, E.S., Wang, H., Durgerian, S., Nikolau, B.J. and Ulrich, T.H. (1993) Plant Physiol. 102, 303-312.

Zhao, Y., Feng, X.H., Watson, J.C., Bottino, P.J. and Kung, S.D. (1994) Plant Molec. Biol. 26, 791-803.

Zimmerman, J.L. (1993) Plant Cell 5, 1411-1423.

What is Claimed is:

- 1. A method of producing apomictic seeds comprising the steps of:
 - (i) transforming plant material with a nucleotide sequence encoding a protein the presence of which in an active form in a cell, or membrane thereof, renders said cell embryogenic,
 - (ii) regenerating the thus transformed material into plants, or carpel-containing parts thereof, and
 - (iii) expressing the sequence in the vicinity of the embryo sac.
- A method according to the preceding claim, wherein the apomictic seeds are of the adventitious embryony type.
- A method according to either of the preceding claims, wherein expression of the sequence yields a protein kinase capable of spanning a plant cell membrane.
- A method according to the preceding claim wherein the kinase is capable of autophosphorylation.
- 5. A method according to any of the preceding claims, wherein the protein is a leucine rich repeat receptor like kinase and comprises a ligand binding domain, a proline box, a transmembrane domain, a kinase domain and a protein binding domain.
- 6. A method according to the preceding claim, wherein the protein lacks a functional ligand binding domain but comprises a proline box, a transmembrane domain, a kinase domain and a protein binding domain.
- A method according to any preceding claim, wherein once incorporated into the cell membrane, the protein binding domain is located intra-cellularly.
- 8. A method according to any preceding claim, wherein the sequence further encodes a cell membrane targeting sequence.

- 9. A method according to any preceding claim, wherein the sequence is that depicted in SEQ ID Nos. 1 or 2 or is complementary to one which hybridizes under stringent conditions with the said sequences and which encodes a membrane bound protein having kinase activity.
- 10. A method according to any preceding claim, wherein the sequence is modified in that known mRNA instability motifs or polyadenylation signals are removed and/or codons which are preferred by the plant into which the sequence is to be inserted are used so that expression of the thus modified sequence in the said plant yields substantially similar protein to that obtained by expression of the unmodified sequence in the organism in which the protein is endogenous.
- 11. A method according to any preceding claim, wherein expression of the sequence is under control of an inducible or developmentally regulated promoter.
- 12. A method according to the preceding claim, wherein expression of the sequence is under control of one of the following: a promoter which regulates expression of SERK genes in planta, the carrot chitinase DcEP3-1 gene promoter, the Arabidopsis AtChitlV gene promoter, the Arabidopsis LTP-1 gene promoter, the Arabidopsis bel-1 gene promoter, the petunia fbp-7 gene promoter, the Arabidopsis ANT gene promoter, the promoter of the O126 gene from Phalaenopsis.
- 13. A method according to any of the preceding claims, wherein the sequence is expressed in the somatic cells of the embryo sac, ovary wall, nucellus, or integuments.
- 14. A method according to any of the preceding claims, wherein the endosperm within the apomictic seed results from fusion of polar nuclei within the embryo sac with a pollenderived male gamete nucleus.
- 15. A method according to the preceding claim, wherein the sequence encoding the protein is expressed prior to fusion of the polar nuclei with the male gamete nucleus.

- 16. DNA comprising a sequence encoding a protein the presence of which in an active form in a cell, or membrane thereof, renders said cell embryogenic.
- 17. DNA according to claim 16, wherein the protein is a leucine rich repeat receptor like kinase and comprises a ligand binding domain, a proline box, a transmembrane domain, a kinase domain and a protein binding domain, the ligand binding domain optionally being absent or functionally inactive.
- 18. DNA according to either of claims 16 or 17 comprising a DNA sequence encoding a N-terminal protein fragment having the following amino acid sequence: Gin Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val Thr Cys Asn.
- 19. DNA according to claim 18 comprising a DNA sequence encoding a N-terminal protein fragment having the following amino acid sequence: Val Xaa Gin Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val Thr Cys Asn with Xaa being a variable amino acid, but preferably Leu or Val.
- 20. DNA according to claim 19 comprising a DNA sequence encoding a N-terminal protein fragment having the following amino acid sequence: Val Xaa Gln Ser Trp Asp Pro Thr Leu Val Asn Pro Cys Thr Trp Phe His Val Thr Cys Asn Xab Xac Xad Xae Val Xaf Arg Val Asp Leu Gly Asn Xag Xah Leu Ser Gly His Leu Xai Pro Glu Leu Gly Xaj Leu Xak Xal Leu Gln with Xaa to Xak being a variable amino acid, but preferably

Xaa = Leu or Val

Xab = Asn or Gin

Xac = Gluor Asportis

Xad = AsnorHis

Xae = Seror Argor Gin

Xaf = le or Thr

Xag = Ala or Ser

Xah=GluorAsn

Xai = Valor Ala

Xaj=ValorLys

Xak=LvsorGlu

- 106 -

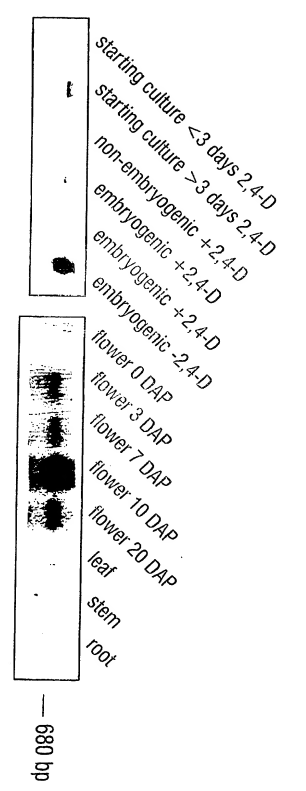
Xal = Asn or His

- 21. DNA comprising a sequence encoding a protein having the sequence depicted in SEQ ID No. 3, or a protein substantially similar thereto which is capable of being membrane bound and which has kinase activity.
- 22. DNA comprising a sequence encoding a protein having the sequence depicted in SEQ ID No. 21, or a protein substantially similar thereto which is capable of being membrane bound and which has kinase activity.
- 23. DNA comprising a sequence encoding a protein having the sequence depicted in SEQ ID No. 33, or a protein substantially similar thereto which is capable of being membrane bound and which has kinase activity.
- 24. DNA comprising a sequence encoding a protein having the sequence depicted in SEQ ID Nos. 23, 25, 27, 29 and 31, or a protein substantially similar thereto which is capable of being membrane bound and which has kinase activity.
- 25. DNA according to any preceding claim, comprising a DNA having the sequence depicted in SEQ ID Nos. 1 or 2 or a sequence which is complementary to one which hybridizes under stringent conditions with the said sequences and which encodes a membrane bound protein having kinase activity.
- 26. DNA according to any preceding claim, comprising a DNA having the sequence depicted in SEQ ID No: 20 or a sequence which is complementary to one which hybridizes under stringent conditions with the said sequences and which encodes a membrane bound protein having kinase activity.
- 27. DNA according to any preceding claim, comprising a DNA having the sequence depicted in SEQ ID No: 32 or a sequence which is complementary to one which hybridizes under stringent conditions with the said sequences and which encodes a membrane bound protein having kinase activity.

- 28. DNA according to any preceding claim, comprising a DNA having the sequence depicted in SEQ ID Nos: 22, 24, 26, 28 and 30 or a sequence which is complementary to one which hybridizes under stringent conditions with the said sequences and which encodes a membrane bound protein having kinase activity.
- 29. DNA according to any of the preceding claims, which further encodes a cell membrane targeting sequence.
- 30. DNA according to any one of the preceding claims, in which the protein encoding region is under expression control of a developmentally regulated or inducible promoter.
- 31. DNA according to claim 30, wherein the promoter is one of the following: a promoter which regulates expression of SERK genes *in planta*, the carrot chitinase DcEP3-1 gene promoter, the *Arabidopsis* AtChitlV gene promoter, the *Arabidopsis* LTP-1 gene promoter, the *Arabidopsis* bel-1 gene promoter, the petunia fbp-7 gene promoter, the *Arabidopsis* ANT gene promoter, the promoter of the O126 gene from *Phalaenopsis*; the *Arabidopsis* DMC1 promoter, the pTA7001 inducible promoter.
- 32. DNA according to any preceding claim, wherein said DNA is a recombinant DNA.
- 33. DNA according to any preceding claim, wherein the sequence is modified in that known mRNA instability motifs or polyadenylation signals are removed and/or codons which are preferred by the plant into which the DNA is to be inserted are used so that expression of the thus modified DNA in the said plant yields substantially similar protein to that obtained by expression of the unmodified DNA in the organism in which the protein is endogenous.
- 34. DNA which is complementary to that which hybridizes under stringent conditions with the DNA of any one of claims 16 to 29.
- 35. A vector containing a DNA sequence as claimed in any one of claims 16 to 34.
- 36. Plant cell transformed with the DNA of any one of claims 16 to 34 or the vector of claim 35, which contains the DNA stably incorporated into its genome.

- 37. Plant cell according to claim 36, which is part of a whole plant.
- 38. Plants transformed with the DNA of any one of claims 16 to 34 or the vector of claim 35, the progeny of such plants which contain the DNA stably incorporated, and/or the apomictic seeds of such plants or such progeny.
- 39. Plants transformed with the DNA comprised by the recombinant DNA of claims 16 to 34.
- 40. Use of the DNA of any one of claims 16-34 in the manufacture of apomictic seeds.
- 41. Plants which are derived from apomictic seeds obtainable by the method of any one of claims 1-15 or 40.
- 42. A method of obtaining cultivars comprising the steps of fertilizing plants with the pollen of the plants of either of claims 38, 39 or 40 and cultivars which result from the said method.
- 43. A method of obtaining embryogenic cells in plant material, comprising transforming the material with a recombinant DNA sequence as claimed in any one of claims 16-34, the DNA comprised by the recombinant DNA of any one of claims 16 to 34, or the vector of claim 35, expressing the sequence in the material or derivatives thereof and subjecting the said material or derivatives to a compound which acts as a ligand for the gene product of the said sequence.
- 44. A method according to the preceding claim, wherein the sequence encodes a leucine rich repeat receptor like kinase, and the compound is a phyto-hormone.
- 45. A method of generating somatic embryos under *in vitr*o conditions wherein the SERK protein is overexpressed ectopically.
- 46. A bag containing apomictic seeds obtainable by the method of any one of claims 1-15 or 40.

Hig. 1



SUBSTITUTE SHEET (RULE 26)

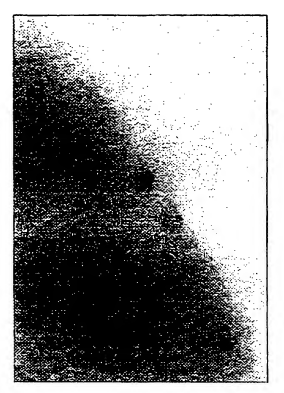


Fig. 2

FIG. 2A

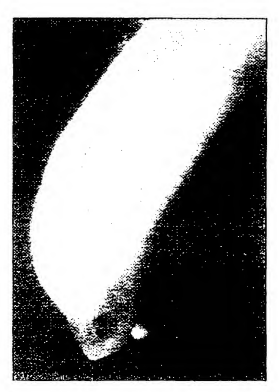


FIG. 2B

SUBSTITUTE SHEET (RULE 26)

Hig. 3

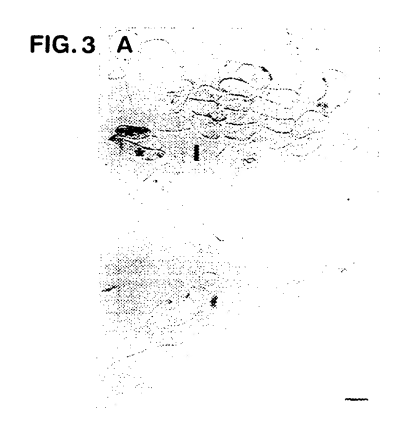


FIG.3 B

FIG.3 C

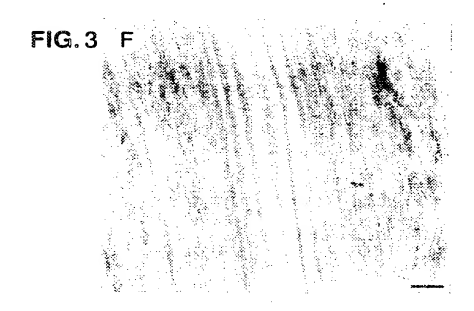
eg

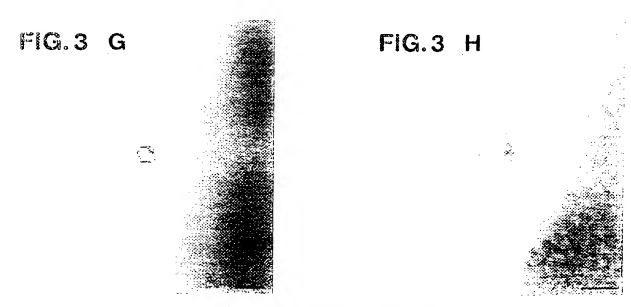
SUBSTITUTE SHEET (RULE 26)

Hin: 3 (cont.)



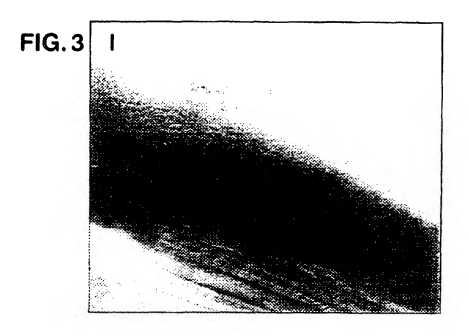


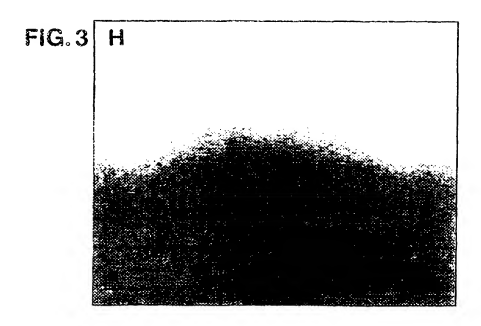




SUBSTITUTE SHEET (RULE 26)

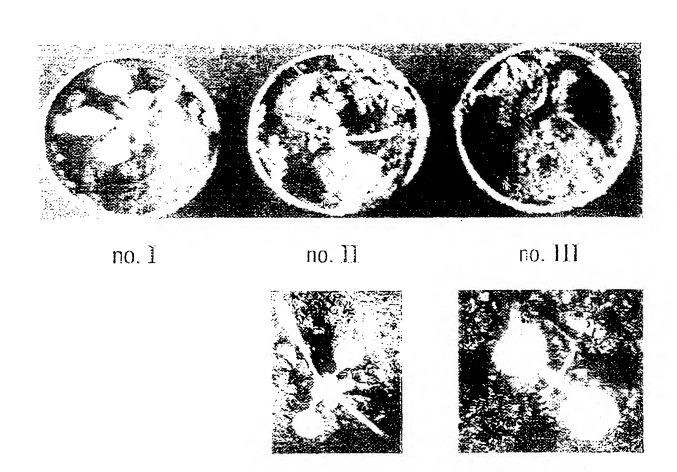
Hig: 3 (cont.)

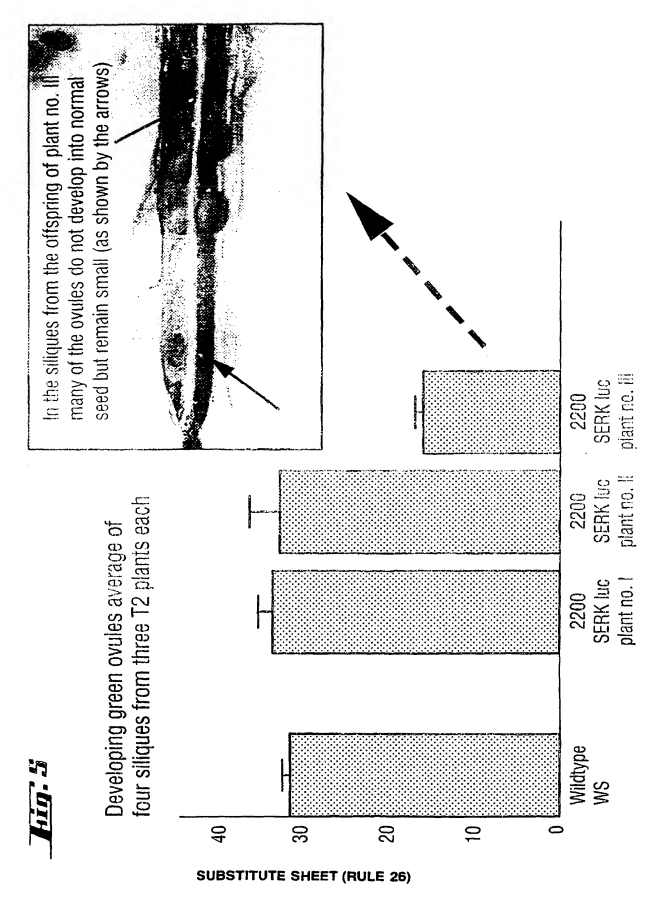




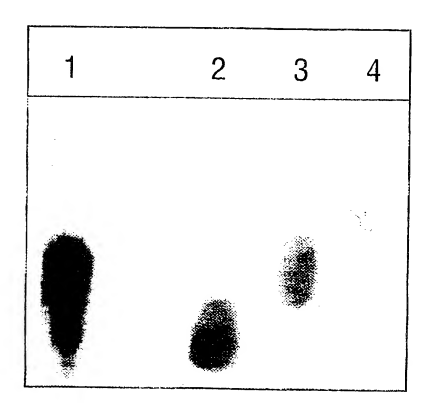
SUBSTITUTE SHEET (RULE 26)

Hig. 4





<u> Hig: 6</u>



SUBSTITUTE SHEET (RULE 26)

A. CLASS IPC 6	IFICATION OF SUBJECT MATTER C12N15/82 C12N9/12 C07K14/	/415 A01H5/00		
According	to International Patent Classification (IPC) or to both national clas	sification and IPC		
	SEARCHED			
Minimum d IPC 6	locumentation searched (classification system followed by classific C12N A01H C07K	ation symbols)		
	tion searched other than minimum documentation to the extent tha			
	lata base consulted during the international search (name of data b	ase and, where practical, search terms used)	
	IENTS CONSIDERED TO BE RELEVANT		<u> </u>	
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.	
х	HANNA, W.W. AND BASHAW, E.C.: "ITS IDENTIFICATION AND USE IN PLBREEDING" CROP SCIENCE, vol. 27, November 1987, pages 1136-1139, XP002040859 see page 1138, left-hand column	1,2		
		-/		
	ner documents are listed in the continuation of box C.	Patent family members are listed	in annex.	
A document defining the general state of the art which is not considered to be of particular relevance *E* earlier document but published on or after the international filing date *L* document which may throw doubts on priority claim(s) or which is cated to establish the publication date of another citation or other special reason (as specified) *O* document referring to an oral disclosure, use, exhibition or other means *P* document published prior to the international filing date but later than the priority date claimed Date of the actual completion of the international search		"T' later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention." "X' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is taken alone. "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family Date of mailing of the international search report 3 0. 09. 97		
Name and m	nailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rigiwijk Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl, Faxc (+ 31-70) 340-3016	Authorized officer Holtorf, S		

Form PCT/ISA/210 (second sheet) (July 1992)

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Category	Chauton of document, with indication, where appropriate, of the relevant passages	
X	KOLTUNOW, A.M., ETAL .: "APOMIXIS: MOLECULAR STRATEGIES FOR THE GENERATION OF GENETICALLY IDENTICAL SEEDS WITHOUT FERTILIZATION" PLANT PHYSIOLOGY, vol. 108, 1995,	1,2,16, 35-41
Y	pages 1345-1352, XP002040860 page 1345, right column, line 1-7; page 1347, right column; page 1348, right column, line 35-50; page 1349, 1350, 1351, left column, line 18-21; Fig. 2 + 3	9
x	KOLTUNOW, A.M.: "APOMIXIS: EMBRYO SACS AND EMBRYOS FORMED WITHOUT MEIOSIS OR FERTILIZATION IN OVULES" THE PLANT CELL, vol. 5, October 1993, pages 1425-1437, XP002040861	1,2,16, 35-41
Υ	see the whole document	9
X	WO 89 00810 A (MAXELL HYBRIDS INC) 9	1,2,16, 35-41
Y	February 1989 pages 7,15,29-35	9
X	MARTIN, G.B.M., ET AL.: "A MEMBER OF THE TOMATO Pto GENE FAMILY CONFERS SENSITIVITY TO FENTHION RESULTING IN RAPID CELL DEATH" THE PLANT CELL, vol. 6, 1994, pages 1543-1552, XP002040862	21-28, 30,32, 34-39
Y	page 1546; Fig.4	9
x	SONG, W-Y, ET AL .: "A RECEPTOR KINASE-LIKE PROTEIN ENCODED BY THE RICE DISEASE RESISTANCE GENE , Xa21" SCIENCE, vol. 270, 15 December 1995, pages 1804-1806, XP002040863 see the whole document	24,30, 32,35-39
X	NEWMAN, T., ET AL .: "GENES GALORE: A SUMMARY OF METHODS FOR ACCESSING RESULTS FROM LARGE-SCALE PARTIAL SEQUENCING OF ANONYMOUS ARABIDOPSIS cDNA CLONES" PLANT PHYSIOLOGY, vol. 106, 1994, pages 1241-1255, XP000571449 see the whole document	27,28
	-/	

Form PCT/ISA/218 (continuation of second sheet) (July 1992)

		PC1/EP 97/02443
<u> </u>	auon) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	NEWMAN,T., ET AL.: "GENES GALORE: A SUMMARY OF METHODS FOR ASSESSING RESULTS FROM LARGE-SCALE PARTIAL SEQEUNCING OF ANONYMOUS ARABIDOPSIS cDNA CLONES" EMBL SEQUENCE DATA LIBRARY, 30 August 1993, HEIDELBERG, GERMANY, XP002040864 ACCESSION No. T04109	27,28
X	NEWMAN, T., ET AL .: "GENES GALORE: A SUMMARY OF METHODS FOR ASSESSING RESULTS FROM LARGE-SCALE PARTIAL SEQUENCING OF ANONYMOUS ARABIDOPSIS cDNA CLONES" EMBL SEQUENCE DATA LIBRARY, 27 July 1995, HEIDELBERG, GERMANY, XP002040865 ACCESSION No. H37195	. 28
X	NEWMAN, T., ET AL.: "GENES GALORE: A SUMMARY OF METHODS FOR ASSESSING RESULTS FROM LARGE-SCALE PARTIAL SEQUENCING OF ANONYMOUS ARABIDOPSIS cDNA CLONES" EMBL SEQUENCE DATA LIBRARY, 27 July 1995, HEIDELBERG, GERMANY, XP002040866 ACCESSION No. H36800	28
X	DESPREZ, T., ET AL.: "THE ARABIDOPSIS THALIANA TRANSCRIBED GENOME: THE GDR cDNA PROGRAM" EMBL SEQUENCE DATA LIBRARY, 25 June 1994, HEIDELBERG, GERMANY, XP002040867 ACCESSION No. Z34606	28
A	DECROOCQ-FERRANT, V., ET AL.: "PETUNIA HYBRIDA HOMOLOGUES OF shaggy/zeste-white 3 EXPRESSED IN FEMALE AND MALE REPRODUCTIVE ORGANS" THE PLANT JOURNAL, vol. 7, no. 6, 1995, pages 897-911, XP002040868 see page 907, right-hand column	1-46
P,X	SCHMIDT, E.D.L., ET AL.: "A LEUCINE-RICH REPEAT CONTAINING RECEPTOR-LIKE KINASE MARKS SOMATIC PLANT CELLS COMPETENT TO FORM EMBRYOS" DEVELOPMENT, vol. 124, no. 10, 1997, pages 2049-2062, XP002040869 see the whole document	16-18, 21-32, 34-37, 39,45
	-/	

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

	ion) DOCUMENTS CONSIDERED TO BE RELEVANT	Relevant to claim No.
ategory "	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim .vo.
Р,Х	WO 97 11167 A (US GOVERNMENT) 27 March 1997 page 1,5,21,24	1,2,41
>,х	WO 97 10704 A (US GOVERNMENT) 27 March 1997	1,2,41
	see the whole document	İ
İ		
:		

INTERNATIONAL SEARCH REPORT

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 8900810 A	09-02-89	AU 629796 B AU 2255288 A EP 0329736 A CN 1040123 A JP 5503836 T	15-10-92 01-03-89 30-08-89 07-03-90 24-06-93
WO 9711167 A	27-03-97	AU 7367396 A	09-04-97
WO 9710704 A	27-03-97	AU 7367496 A	09-04-97

Form PCT/ISA/210 (patent family annex) (July 1992)

			••
• 7			